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Experimental simulation of magma-carbonate interaction beneath Mt. Vesuvius, Italy

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Abstract

We simulated the process of magma-carbonate interaction beneath Mt. Vesuvius in short duration piston cylinder experiments under controlled magmatic conditions (from 0 to 300 s at 0.5 GPa and 1200 °C), using a Vesuvius shoshonite composition and upper crustal limestone and dolostone as starting materials. Backscattered electron images and chemical analysis (major and trace elements and Sr isotopes) of sequential experimental products allow us to identify the textural and chemical evolution of carbonated products during the assimilation process. We demonstrate that melt-carbonate interaction can be extremely fast (minutes) and results in dynamic contamination of the host melt with respect to Ca, Mg and ⁸⁷Sr/⁸⁶Sr, coupled with intense CO₂ vesiculation at the melt-carbonate interface. Binary mixing between carbonate and uncontaminated melt cannot explain the geochemical variations of the experimental charges in full and convection and diffusion likely also operated in the charges. Physical mixing and mingling driven by exsolving volatiles, seems to be a key process to promote melt homogenisation. Our results reinforce hypotheses that magma-carbonate interaction is a relevant and ongoing process at Mt. Vesuvius and one that may operate not only on a geological, but on a human timescale.

Keywords: Mt. Vesuvius, magma-carbonate interaction, crustal assimilation, CO₂ liberation, experimental petrology

1. Introduction

The Mt. Somma-Vesuvius complex (1281 m a. s. l.) comprises Mt. Somma, the > 25 to 14 kyr strato-volcano, and Mt. Vesuvius, the recent cone which has grown within the older Mt. Somma caldera (e.g., [Rolandi et al. 2004](#); [Peccerillo, 2005](#); [Piochi et al. 2006](#); [Di Renzo et al. 2007](#)) and which we collectively term the “Vesuvius volcanic system” (VVS). The VVS is characterised by variable types of eruption, ranging from effusive lava emission to explosive strombolian, sub-plinian and plinian events ([Orsi et al. 1996](#); [Peccerillo 2005](#); [Paone 2006](#); [Piochi et al. 2006](#)). The VVS is sited on a sequence of Mesozoic and Cenozoic carbonates overlain by the Miocene siltstone sediments of the Campanian Plain ([Brocchini et al. 2001](#); [Civetta et al. 2004](#); [Del Pezzo et al. 2006](#)). Seismic tomography studies have suggested that the top of the Mesozoic carbonate basement lies at around 2.5 - 3 km depth underneath the volcano ([Zollo et al. 2002](#)). The level at ~ 8 - 11 km depth has also been identified as a discontinuity, likely representing the top of the present day magma reservoir, which is probably linked to the base of the Mesozoic carbonate sequence at ~ 10 - 11 km depth ([Zollo et al. 1996](#); [Berrino et al. 1998](#); [Auger et al. 2001](#)). The VVS has been the focus of attention ever since the famous AD 79 eruption, being one of the most hazardous volcanic systems in Europe due to the densely populated Neapolitan region in its shadow. Although much is known concerning the volcano and its internal workings, a comprehensive understanding of the role of magma-crust interaction processes at the VVS has not yet been achieved.

The VVS produces slightly silica-undersaturated K-basalts and K-trachytes to highly silica-undersaturated K-tephrites and K-phonolites ([D’Antonio et al. 1999](#); [Paone 2006](#); [Piochi et al. 2006](#)). The compositional variability of these magmas has been attributed to: *i*) mantle source variability (e.g., [Ayuso et al. 1998](#); [Peccerillo 1999](#); [Somma et al. 2001](#)), *ii*) differentiation and magma mixing (e.g., [Turi and Taylor 1976](#); [Civetta et al. 1991](#)) and *iii*) contamination through carbonate assimilation (e.g., [Rittmann, 1933](#); [Fulignati et al. 1995](#); [Del Moro et al. 2001](#); [Gilg et al. 2001](#)). The role of carbonate assimilation has recently been documented for a

significant number of volcanic systems emplaced in carbonate-rich crust, e.g., Popocatepetl, Colli Albani, and Merapi (e.g., Goff et al. 2001; Dallai et al. 2004; Schaaf et al. 2005; Chadwick et al. 2007; Freda et al. 2008). In fact, carbonate assimilation has been shown to occur over short timescales (minutes in experiments) and thus the effect of carbonate decomposition in natural systems is thought to release significant amounts of CO₂ on timescales of hours to days (Deegan et al., 2010, 2011; Troll et al., 2012a). Such CO₂ influx into a volcanic system may not only have the potential to modify magma composition but also the style of eruptive activity (Deegan et al. 2010, 2011; Dallai et al. 2011; Freda et al. 2011; Troll et al. 2012a, b). At the VVS, the presence of a thick carbonate basement, the frequent occurrence of skarn xenoliths, and the anomalous gas chemical compositions unequivocally suggest that contamination through carbonate assimilation is a relevant process and may play an important role in the volcano's magma evolution and its eruptive behaviour and styles (cf. Freda et al. 1997; Del Moro et al. 2001; Gilg et al. 2001; Fulignati et al. 2004, 2005; Iacono-Marziano et al. 2009; Dallai et al. 2011). In this study, we report on a set of experiments designed to examine magma-carbonate interaction processes under controlled conditions (0.5 GPa and 1200 °C), employing a composition representative of primitive VVS magmas and various local carbonates. Previously, magma-carbonate interaction experiments were run over hours to days to establish mineral phase relationships and the liquid line of descent in magmatic products affected by carbonates (e.g., Freda et al., 2008; Iacono Marziano et al. 2008; Mollo et al. 2010). In contrast, our experimental series is designed to perform very short duration experiments (0, 60, 90 and 300 s; cf. Deegan et al. 2010), allowing the preservation of textures and phases developed during progressive carbonate break-down and degassing. Coupled with *in-situ* elemental and Sr-isotope constraints on our experiments, our data provide a more comprehensive understanding of the dynamics of magma-carbonate interaction in the VVS.

2. Experimental methods

The experiments were conducted in an end-loaded piston-cylinder apparatus at the HP-HT Laboratory of Experimental Volcanology and Geophysics of Istituto Nazionale di Geofisica e Vulcanologia (INGV, Rome, Italy). Experiments were carried out at 0.5 GPa equivalent to ca. ~ 13 - 14 km depth. This is the lowest pressure to which this end-loaded piston-cylinder apparatus is calibrated for, but is similar to the pressure inferred for the VVS magma reservoir (≥ 10 km depth; Zollo et al. 1996, 1998; Auger et al., 2001). The experimental temperature of 1200 °C was calculated as the liquidus temperature of the starting composition at 0.5 GPa and initial water content of 2 wt. %, using the MELTS algorithm (Ghiorso et al. 1994). The experimental temperature was reached in 6 minutes, which is sufficiently fast to preserve the textural relationship of the carbonate phase in the shortest duration experiments and allow inspection of the interaction between carbonate and the resulting melts in detail (see Deegan et al. 2010). Our experimental approach thus approximates the likely physical and chemical conditions of direct magma-carbonate interaction in the VVS.

The starting materials used in our experiments were a shoshonitic lava flow from the VVS (Di Renzo et al. 2007) and limestone and dolostone from the local Procida carbonate formation (e.g., Barberi et al. 1981; Civetta et al. 1991). The shoshonitic lava flow represents one of the least evolved rocks outcropping in the Neapolitan area. A shoshonite sample was first crushed, powdered, and doped with 2 wt. % H₂O (added by microsyringe into an experimental capsule) and then melted at 1250 °C and 0.5 GPa to produce a hydrated shoshonite glass. The hydrated glass was analysed for its major element composition (see **Table 1**) and was verified to be free of crystals and crystallites. Then, the hydrated shoshonite glass was powdered and used as the magmatic starting material in our experimental series.

The hydrated shoshonite powder was placed in platinum capsules (3.0 mm diameter, 12.0 mm long) together with fragments of ~ 6 - 8 mg (≤ 3.0 mm side length) of either limestone or dolostone (**Table 1**). The capsules were then welded shut and positioned in tandem into a 19.1 mm NaCl-crushable alumina-pyrex assembly. This means that the two capsules were placed

side by side in the same assembly, one containing the hydrated shoshonite glass and limestone and the other the hydrated shoshonite glass and dolostone. The tandem approach has the advantage that it allows a direct comparison between limestone- and dolostone-bearing experiments as physical parameters (pressure, temperature and time) are otherwise identical. The capsules were surrounded by pyrophyllite powder to prevent water loss (see [Freda et al., 2001](#)) and further details of the experimental approach can be found in [Freda et al. \(2008\)](#) and [Deegan et al. \(2010\)](#).

3. Analytical Methods

The experimental products were analysed using a JEOL-JXA8200 electron microprobe (EMPA) and a JEOL-6500F field emission scanning electron microscope (FE-SEM) at INGV Rome. Microprobe analyses were performed using an accelerating voltage of 15 kV, a beam current of 5 nA, and a beam diameter of 5 μm for glass and 1 μm for mineral analyses (see, e.g., [Iezzi et al., 2008](#), for analytical details). Sodium and potassium were analysed first to reduce possible volatilisation effects. The average standard deviation (1σ) of major element oxides (in wt. %) is 0.34 for SiO_2 , 0.06 for TiO_2 , 0.11 for Al_2O_3 , 0.10 for FeO , 0.03 for MnO , 0.07 for MgO , 0.16 for CaO , 0.05 for Na_2O , 0.03 for K_2O , 0.04 for P_2O_5 .

Micro-sampling of the experimental products for their Sr isotope ratios was performed at the Arthur Holmes Isotope Laboratory, Department of Earth Sciences, Durham University, UK using a New Wave MicromillTM and following the technique outlined in [Charlier et al. \(2006\)](#). The Micromill consists of a binocular microscope, a mill, a stage, which controls movement reproducible to $\pm 1 \mu\text{m}$, and a computer workstation which integrates all the components ([Charlier et al. 2006](#)). Sampling sites were selected using BSE images to precisely locate the drill points and avoid fractures and bubbles. The samples were milled along selected areas to a depth of 30 μm . Milling was carried out under a single drop of Milli-Q water to collect the drilled sample dust. The sample and Milli-Q water mixture was pipetted off and transferred directly to a ‘gold boat’ for processing and micro-Sr column chemistry as described in

Charlier et al. (2006). After processing, the samples were analysed by TIMS using a Thermo-Finnigan Triton system (see Font et al. 2008). During the analysis period (July to August 2010) the international Sr standard NBS-987 was analysed with sample sizes ranging from 3 ng to 600 ng. The overall average Sr isotopic ratios for the NBS-987 measurements during this period ($n = 26$) was 0.710261 ± 0.000016 , within error of the published $^{87}\text{Sr}/^{86}\text{Sr}$ value of 0.710248 ± 0.000023 (2σ) ($n = 427$) reported by Thirlwall et al. (1991). Aliquots of the dissolved milled samples were analysed for their trace element concentrations by inductively coupled plasma mass spectrometry (ICP-MS) using a Thermo Electron Element II system at Durham University, UK. Procedural details are provided in Font et al. (2008). Total procedural blanks ($n = 5$) were less than 22 pg for all elements analysed. Sr blanks averaged $0 \pm 1 \text{ pg}$ (2σ , $n = 5$).

4. Results

4.1. Features of the experimental products

The experimental conditions for all ten experiments are given in **Table 2** and representative electron microprobe analytical traverses of two experimental products are reported in **Table 3**. Three major phases were identified: *i*) two distinct glasses (representing melts), Ca-normal and either Ca- or Ca-Mg-rich, which are identified in BSE images as dark and bright in colour, respectively. A contamination front (CF) usually separates the two glass domains (**Fig. 1**). *ii*) Crystalline phases, such as calcite, pyroxene and olivine (**Figs. 2 and 3**), and *iii*) a vapour phase represented by bubbles (e.g., **Fig. 1**). In the following section the main features of the experimental products are described to explain the dynamics of magma-crust interaction in the experiments and to report on the differences between limestone- and dolostone-bearing experiments.

Limestone-bearing experiments

In the limestone-bearing experimental products, for $t_d = 0$ to 90 s, two distinct glass regions were observed (**Fig. 1a-c**): *i*) a Ca-normal glass domain, very similar in composition to the shoshonite starting material (av. 49.85 wt. % SiO_2 , 9.98 wt. % CaO) and *ii*) a Ca-rich glass domain with an interface between the two (CF). Within the CF region, the Ca-normal glass domain shows progressive depletion in silica and enrichment in CaO , grading into the Ca-rich melt domain (**Fig. 4**), with aluminium strongly mimicking the behaviour of silica. The interfacial regions (CF) hence involve progressive physico-chemical mixing between the two dominant glass domains and an interfacial boundary between the Ca-rich and Ca-normal domains is observed in all experiments. The solid phases are dendritic micro-crystals of calcite (**Fig. 2**) produced in Ca-rich areas of the Ca-rich glass domains. These crystals represent quench crystals formed from local enrichment of former calcite components in the experimental melts. Notably, the $t_d = 300$ s run is characterised by a more homogeneous Ca-rich glass, vapour (bubbles) and no crystals (**Fig. 1d**).

Dolostone-bearing experiments

The dolostone-bearing experimental products contain either glass and vapour phases ($t_d = 0$ s; **Fig. 1e**) or crystals, glass and vapour phases ($t_d = 60$ to 300 s; **Fig. 1f-h**). The results of chemical traverses through the various glass regions are shown in **Figure 5**. The dolostone-bearing experiments show a normal glass with chemical composition similar to the starting material, a contaminated glass (Ca-Mg enriched), and CF regions in between the two. In the CF areas, it is possible to distinguish a progressive dilution of silica and coeval calcium and magnesium enrichment relative to the shoshonite starting material (**Fig. 5**). Crystalline phases in the dolostone-bearing experiments are ferromagnesian olivine and pyroxene. Analyses of all mineral phases are provided in the Online Resource 1 (**Table OR1**). The ternary diagram in **Figure 3** shows the crystalline phases in the Ca_2SiO_4 - Mg_2SiO_4 - Fe_2SiO_4 and in the CaSiO_3 - MgSiO_3 - FeSiO_3 systems. Pyroxene is dominant over olivine and is highly enriched in CaO (> 20 wt %) and MgO (10.01 - 13.16 wt. %), in agreement with its occurrence in the

interfacial area, i.e., depletion in SiO₂ and enrichment in CaO, MgO and Al₂O₃ (**Fig. 3a**). The olivine compositions are close to forsterite (Fo_{95-97.5}), but with a notable enrichment in CaO (1.31 to 2.36 wt %; **Fig. 3b**) and thin Fe-rich, likely more outer rims. The olivine crystals (~10 - 20 µm size) are euhedral and the pyroxene display euhedral to subhedral shapes, which points to direct growth of olivine and pyroxene from contaminated melts (**Fig. 3**).

Vapour phase in experimental products

The vapour phase in our experiments is preserved as numerous vesicles (bubbles) that formed at the contamination front predominantly from exsolved CO₂ (see below). Vesicles were observed in all experimental runs in different quantities, sizes and spatial distributions. For dolostone-bearing experiments, the vesicles' spatial distribution is not random, rather the bubbles tend to concentrate close to contaminated areas. In contrast, for limestone-bearing experiments the bubbles have a tendency to nucleate in the Ca-rich melt domains and migrate into the Ca-normal glass as reflected by different generations of vesicles and micro-bubble fronts (**Fig. 1**). Bubble sizes increase from a micrometer to a millimetre scale with progressive experimental run-time (Blythe et al. 2012), reflecting initial bubble nucleation that is followed by progressive bubble growth and coalescence.

4.2. Strontium isotope compositions in the experimental products

The Sr isotope variations in limestone- and dolostone-bearing experiments are illustrated in **Figures 6** and **7** and the data are reported in **Table 4**. All trace element concentrations in the drilled samples can be found in the Online Resource 2 (**Table OR2**). The micro-Sr-isotope analyses show significant ⁸⁷Sr/⁸⁶Sr variability in each experiment (0.706729±10 to 0.707023±33 in limestone-bearing and 0.707017±8 to 0.707743±10 in dolostone-bearing experiments). The analysed Ca-normal glass has low ⁸⁷Sr/⁸⁶Sr values (0.706729±10 to 0.706896±40) similar to those of the shoshonite starting material (0.706661±8; Di Renzo et al. 2007). In contrast, the more radiogenic ⁸⁷Sr/⁸⁶Sr values (0.706951±17 to 0.707743±10) correspond to Ca- or Ca-Mg-rich domains, with ⁸⁷Sr/⁸⁶Sr ratios that trend towards those of the

carbonates of the Campanian region (0.7072 ± 1 to 0.7093 ± 8 ; Civetta et al. 1991; Piochi et al. 2006; Iannace et al. 2011). Three of the microdrilled experiments were analysed in the CF, showing $^{87}\text{Sr}/^{86}\text{Sr}$ signatures that fall in between the respective Ca- or Ca-Mg-rich and Ca-normal glass compositions of these experiments (Figs. 6 and 7).

5. Discussion

5.1. Effects of magma-carbonate interaction

Magma-carbonate interaction and consequent carbonate assimilation allows for several possible reaction routes, ranging from direct dissolution to formation of a series of intermediate products, such as the various skarn assemblages frequently observed (e.g., Gaeta et al. 2009; Mollo et al. 2010; Troll et al. 2012b). The net result of this interaction is an overall release of CaO by limestone and CaO and MgO by dolostone into the host magmas(s), and associated liberation of CO_2 (cf. Mollo et al. 2012; Troll et al. 2012a). The experiments simulate direct magma-carbonate interaction, which is demonstrated by the chemical profiles and by the abundant occurrence of bubbles in the experimental products. A recent experimental investigation using a basaltic-andesite composition from Merapi volcano and Indonesian carbonate as starting material, and employing analogous pressure (0.5 GPa), temperature (1200 °C) and water content (~ 2 wt. %), showed similar processes of magma-carbonate interaction at work (Deegan et al. 2010). These authors established that carbonate break-down began during the experimental heating phase (approximately 6 min.) and the time required for complete carbonate assimilation was probably no more than ca. 330 s at their target experimental conditions. In contrast to the Merapi experiments, we observe that no original carbonate is preserved in any of our experimental products (Figs. 1 and 2), implying that carbonate dissociation and complete dissolution occurred before the target temperature of 1200 °C was reached. The highest degree of physico-chemical mixing between melt domains occurs between $t_d = 60$ and 90 s (i.e., mixing initially increases with time; Figs. 8 and 9). At $t_d \sim 300$ s, contaminated melt dominates almost the entire volume of the experimental charge,

leaving normal melts at the rims only, i.e., mixing is approaching completion. It should be noted though that calcite (or alternatively pyroxene and olivine) crystallised where local enrichment of certain elements was the result of magma-carbonate interaction and implies that chemical homogenisation between melts is limited within the timeframe of our experiments, i.e., the run-time for the experiments does not permit complete homogenisation. The Vesuvius experiments, in contrast to the Merapi ones, therefore display a significantly shorter timescale of magma-carbonate interaction, suggesting faster reaction rates between the shoshonite melt and the carbonate (limestone and dolostone) under otherwise similar pressure and temperature conditions.

The chemical variations between glass domains in limestone- and dolostone-bearing experiments can be approximated by binary mixing models with the starting materials (limestone, dolostone and shoshonite) as end-members. Based on these models, we provide an estimate of the maximum degree of mixing between melts in each experimental step. The degree of mixing varies mainly with the experimental duration (t_d) (**Figs. 8 and 9**) and maximum mixing degrees are often confined to spatially limited zones of about 100-200 μm in experiments up to 90 s (**Figs. 4 and 5**).

The limestone-bearing experiments reflect this variation on the maximum degree of mixing between carbonate and shoshonite through time (**Fig. 8**; see **Online Resource 3 Fig OR1**) and similar degrees of mixing are observed for elements such as MgO , Al_2O_3 or NaO_2 . The maximum degree of limestone component in the melt phase (limestone: shoshonite, weight ratio) is *i*) ~ 40:60 at $t_d = 0$ s, *ii*) ~ 50:50 at $t_d = 60$ s, and *iii*) ~ 60:40 at $t_d = 90$ s. At *iv*) $t_d = 300$ s, the resultant melt is almost homogenous, representing a mixture between dissolved carbonate and shoshonite melt of about 25:75. For the dolostone-bearing experiments, the maximum degree of dolostone component in the melt phase (dolostone: shoshonite, weight ratio) for experimental duration is (**Fig. 9**; see **Online Resource 3 Fig OR2**): *i*) ~ 15:85 at $t_d = 0$ s, *ii*) ~ 45:55 at $t_d = 60$ s, *iii*) ~ 45:55 at $t_d = 90$ s, and *iv*) about 25:75 at the longest

experimental duration of 300 s. These models demonstrate that the degree of mixing is time-
 dependent and generally greater for the longer duration experiments. For the experiments
 between 0 and 90 s, the resultant melts are inhomogeneous and in disequilibrium since the
 system undergoes various degrees of diffusive and convective mixing in addition to crystal
 growth and resorption (e.g., calcite, pyroxene and olivine). In contrast, at $t_d = 300$ s, in both
 limestone- and dolostone-bearing experiments, a hybrid melt containing about 25 wt. % CaO
 begins to form as the result of pervasive mixing and progressive homogenisation (**Figs. 8 and**
9; Online Resource 1). Pure binary mixing between two melt end-members should follow a
 linear relationship for a regular element pair (e.g., CaO vs. SiO_2), which, however, is not
 always observed in our experiments (**Figs. 8 and 9**). The observed variations may reflect
 different mobility of trace and major elements over very short timescales during incipient
 mingling and mixing, producing complex and fluctuating patterns that deviate from straight
 mixing trends. This deviation is most pronounced in the dolostone-bearing experiments,
 however, where the mixing trajectory appears to move furthest from the straight mixing trend,
 probably a result of crystal growth (olivine and pyroxene) and associated Mg removal from
 the melt that will shift the Ca/Mg ratio of the melt away from that of an ideal mixture ([De](#)
[Campos et al. 2008; Perugini et al. 2006, 2008](#)). We also estimated “apparent diffusivities” of
 SiO_2 , Al_2O_3 , MgO and CaO across the contamination front in the limestone-bearing
 experiments (see **Online Resource 4**). The results yield apparent diffusivities in m^2/s of
 $8.5 \cdot 10^{-10}$ to $8.6 \cdot 10^{-7}$ for SiO_2 , $8.5 \cdot 10^{-10}$ to $1.9 \cdot 10^{-9}$ for Al_2O_3 , $8.5 \cdot 10^{-10}$ to $1.1 \cdot 10^{-8}$ for MgO
 and $3.4 \cdot 10^{-11}$ to $6.3 \cdot 10^{-6}$ for CaO in the 60 and 300 s experiments, respectively (**Online**
Resource 4). The determined values are between two and five orders of magnitude greater
 than well-established diffusivities for these elements (e.g., [Watson 1982; Watson and](#)
[Jurewicz 1984; Baker 1991; Lesher et al. 1996; Liang et al 1996; Zhang 1993 and 2010](#)) and,
 moreover, they vary between the different duration experiments. These data imply that
 although diffusion is a relevant process at play, the overall elemental gradients observed must

be the result of a combination of processes including mingling, mixing, convection, and local crystal growth (**Figs. 1, 8 and 9**). These additional processes can all affect the melt interface and thus overprint diffusion profiles in the experimental charges. In our experiments, the process of physical mixing and mingling observed between the melt domains is driven by volatiles that exsolve during carbonate assimilation (**Fig. 1**), which likely contributes most strongly to melt homogenisation on the timescale of our experiments. Crystal growth, on the other hand, is influenced by carbonate break-down in our experiments. Limestone, for example, contributes to local calcite-saturation whereas dolostone provides additional CaO and MgO thereby promoting crystallisation of Ca-rich pyroxene and Ca-Mg-rich olivine (cf. [Metz and Milke, 2012](#)). The crystallisation of these mineral phases in our experiments is a function of the state of the contaminated melt, i.e., it is dependent on melt domains that are locally enriched in CaO and MgO prior to full homogenisation. Notably, the growth of high-Mg olivine from dolostone contaminated melt raises the possibility that high-Fo olivine is not always an indicator for primitive or primary magma compositions in volcanic systems and particularly not in the VVS.

5.2. $^{87}\text{Sr}/^{86}\text{Sr}$ composition of the experimental products

The $^{87}\text{Sr}/^{86}\text{Sr}$ isotope variations provide a first order approximation of the degree of carbonate assimilation in our experiments. The Sr isotope ratios and Sr concentration for Mt. Vesuvius shoshonitic magmas range between 0.7067 and 0.7071 and from 735 to 1093 ppm, respectively ([Di Renzo et al. 2007](#)). The $^{87}\text{Sr}/^{86}\text{Sr}$ isotope ratios and Sr concentration of carbonate-rich lithologies from the Campanian region were previously thought to range between 0.7075 and 0.7090 and from 500 to 1000 ppm, respectively ([Civetta et al. 1991](#); [Piochi et al. 2006](#); [Del Moro et al. 2001](#)). More recent studies on the same carbonates now also report lower values of $^{87}\text{Sr}/^{86}\text{Sr}$ to be present, e.g., 0.7072 to 0.7074 ([Iannace et al. 2011](#)) with associated Sr concentrations of as little as 41 to 151 ppm, implying that these carbonates are highly heterogeneous in their Sr concentrations and $^{87}\text{Sr}/^{86}\text{Sr}$ composition (see below).

318 The Sr-isotope profiles of the experimental products display a broad correlation of higher
 319 $^{87}\text{Sr}/^{86}\text{Sr}$ ratios with increasing CaO wt. %, i.e., with the most contaminated drill samples,
 320 whereas the non-contaminated glasses show Sr isotope ratios that overlap with those of the
 321 natural shoshonite starting material (**Fig. 10a**). The $^{87}\text{Sr}/^{86}\text{Sr}$ ratios observed in the CF regions
 322 usually fall in between these extremes and frequently form mixtures with intermediate
 323 $^{87}\text{Sr}/^{86}\text{Sr}$ signatures (**Figs. 6 and 7**). Indeed, in this respect, the isotopic variations observed
 324 within each individual experiment can be explained as the result of mixing between “normal”
 325 and “contaminated” melts. The situation appears somewhat more complex, however, when
 326 using binary mixing trajectories between the published values for the shoshonite and for the
 327 various carbonate $^{87}\text{Sr}/^{86}\text{Sr}$ values as well as for a high (500-1000 ppm) and a low (<500 ppm)
 328 Sr concentration group that are now available in the literature (e.g., [Civetta et al. 1991](#); [Piochi](#)
 329 [et al. 2006](#); [Del Moro et al. 2001](#); [Di Renzo et al., 2007](#); [Iannace et al. 2011](#)). For the VVS
 330 shoshonite, a $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.7067 and Sr concentration of 735 ppm was chosen (**Table 1**).
 331 For the carbonates, five values were selected (C₁-C₅) because their Sr-isotope ratios are so
 332 variable that our single compositions cannot represent the available data range on limestone
 333 and dolostone from the Campanian region. For example, the high Sr concentration carbonate
 334 (500-1000 ppm) cannot explain the range of Sr isotopic values measured in the experiments as
 335 only one drill sample, the most radiogenic one (D-V2.4, 0.707743), falls into the shoshonite –
 336 C₁– C₂ – mixing space (**Fig. 10b**). The best fit for the remaining drill data is achieved using
 337 low Sr (<500 ppm) carbonates (C₂-C₅). Employing these possible end-member(s), all
 338 remaining experimental data fall into the theoretically possible mixing space. This observed
 339 range of $^{87}\text{Sr}/^{86}\text{Sr}$ in the experimental Ca-rich melts implies that on the one hand the
 340 contaminant used in our experiments is highly inhomogeneous on a mm to sub-mm-scale,
 341 which is consistent with the realisation that the Campanian carbonates are heterogeneous
 342 regarding their Sr-isotope values as well as their Sr-concentrations (e.g., [Iacono-Marziano et](#)
 343 [al. 2008](#); [Iannace et al. 2011](#)). On the other hand, most carbonate material in our experiments

appears to have lower Sr concentrations than the shoshonite melt (see **Table 1**; [Di Renzo et al., 2007](#)). A contaminated shoshonite may therefore be less radiogenic than previous predictions of contamination had suggested on the basis of the high $^{87}\text{Sr}/^{86}\text{Sr}$ and Sr concentration values available for the Procida formation at that time (e.g., [Ayuso et al. 1998](#); [Iacono-Marziano et al. 2008](#)). Considering the full range of possible mathematical solutions, including the new low Sr and low $^{87}\text{Sr}/^{86}\text{Sr}$ end-members ([Iannace et al. 2011](#); **Table 1**), the most intensely contaminated experimental drill-samples then calculate to mixtures of up to between 55 and 75% carbonate-derived $^{87}\text{Sr}/^{86}\text{Sr}$ relative to unmodified shoshonite (**Fig. 10b**). The noteworthy aspect of these results is that relatively primitive compositions at the VVS (e.g., shoshonite) may represent contaminated magma even in the absence of particularly high Sr-isotope ratios or high Sr elemental concentrations, as large amounts of contamination by a low $^{87}\text{Sr}/^{86}\text{Sr}$ and low Sr ppm carbonate will only exhibit a small effect on the Sr isotope composition of the magma. A CaO and Sr relationship, as previously predicted for substantial limestone assimilation, cannot be expected if low-Sr isotope contaminants and variable Ca and Mg ratios in limestones versus dolostones (and their resulting calc-silicate skarns) are considered to influence the system in addition to pure limestone uptake. The lack of a clear CaO-Sr relationship is hence not an argument against carbonate assimilation (cf. [Ayuso et al. 1998](#)). Coupled with the occurrence of high-Mg olivine and pyroxene in our dolostone-bearing experimental products, a mineral association known to occur in some primitive VVS samples (e.g., [Dallai et al. 2011](#)), low-Sr dolostone may in fact be a key contaminant for the VVS.

5.3. CO_2 liberation during magma-carbonate interaction

Dissolved volatiles in silicate melts play an important role in defining physical properties of magmas for, e.g., density, viscosity, vesicularity, and thus influence ascent behaviour and eruptive styles (e.g., [Dingwell 1996](#); [Baker et al. 2005](#); [Deegan et al. 2010](#); [Lesne et al. 2010](#)). The most abundant volatiles in magmas are H_2O and CO_2 , which are commonly dissolved in

different percentages depending on melt composition, pressure, temperature and degree of saturation (e.g., [Zhang and Stolper 1991](#)). In particular, solubility experiments on shoshonite under conditions similar to those used in this study, i.e., 1250 °C and 0.4 GPa, show that water solubility is about 8 wt. %, while CO₂ solubility is limited to < 3000 ppm. Under CO₂ saturated conditions (> 3000 ppm), only 2 wt. % H₂O is soluble, however ([Di Matteo et al. 2006](#); [Vetere et al. 2011](#)). This is broadly consistent with H₂O and CO₂ solubilities predicted for shoshonite by the numerical model of [Papale et al. \(2006\)](#), which yields solubilities of 1.9 wt. % and 3449 ppm for H₂O and CO₂, respectively, at the pressure and temperature conditions of our experiments. In our experiments, therefore, H₂O and CO₂ coexist in the melt. During carbonate break-down and assimilation, the progressive increase of CaO and CO₂ in the melt will affect both the CO₂ and H₂O solubilities (see **Table 3**; [Holloway 1976](#); [Watson et al. 1982](#); [Blank and Brooker 1994](#); [Holloway and Blank 1994](#); [Dixon 1997](#); [Botcharnikov et al. 2005](#); [Moore, 2008](#)) and will provide CO₂ to the melt until volatile saturation is achieved. At this point, a fluid phase will form that will likely contain only a small portion of H₂O since the solubility of water in our experimental system is close to our starting content of 2 wt. % according to the models of [Papale et al. \(2006\)](#) and [Vetere et al. \(2011\)](#). Either model implies that although some H₂O is transferred from the melt to the fluid phase on CO₂ saturation, most of the water remains in the melt. Carbonate assimilation will then progressively add more CO₂ to the fluid phase, meaning that in our experimental charges the final mixed H₂O-CO₂ vapour phase is strongly CO₂ dominated.

The potential of CO₂ released by carbonate assimilation in the experimental charges can be assessed using an average amount of carbonate and shoshonitic melt of 6.8 and 29 mg, respectively. The complete break-down of carbonate during our experiments will liberate 2.99 mg of CO₂, which corresponds to 4.89 wt. % of CO₂ in the experimental charge as a whole (versus ≤ 2 wt. % H₂O). Therefore, the experimental approach presented here shows that CO₂

liberation during carbonate assimilation can be significant under crustal magma reservoir conditions and may be an important factor in promoting explosive behaviour at the VVS.

6. Implications of the Vesuvius volcanic system

Magma-carbonate interaction is a relevant process in the VVS and is characterised by: *i*) the abundance of high temperature skarn xenoliths in the erupted products (e.g., [Fulignati et al. 1998, 2004, 2005](#); [Del Moro et al. 2001](#)), *ii*) crustally-derived CO₂ detected via the chemical and isotope composition of Vesuvius fumaroles (e.g., [Iacono-Marziano et al. 2009](#)), and *iii*) the $\delta^{18}\text{O}$ isotope composition of mafic VVS minerals that experienced interaction with sedimentary carbonate already at considerable depths ([Dallai et al. 2011](#)). Judging from our experiments, interaction between carbonate and Vesuvius shoshonite is indeed a viable and likely very fast process (minutes to hours). Moreover, we have shown how this process will progressively enrich the host melt in CaO and likely also MgO (from dolostone), while generating a CO₂-rich fluid phase. The experimental products show that CO₂ is generated directly at the melt-carbonate interface. In nature, this gas phase would form bubbles at crustal levels ([Holloway and Blank 1994](#)) and be transported upwards through the magma plumbing and conduit system. Therefore, CO₂ fluxes at the VVS are probably highly variable over different timescales, and CO₂ emissions measured in volcanic fumaroles at Vesuvius, that show a dominance of crustal over mantle-derived CO₂ (e.g., [Iacono-Marziano et al. 2009](#)), may be explained by the process modeled in our experiments. Conceivably, large quantities of CO₂ may be rapidly liberated during, e.g., renewed magma injection into shallow levels of the VVS reservoir system, where magma-carbonate interaction and carbonate assimilation have the potential to promote erratic explosive behaviour over short timescales (cf. [Deegan et al. 2010, 2011](#); [Freda et al. 2011](#)). This implication would appear to be relevant to other volcanic systems as well, notably Merapi, Indonesia ([Chadwick et al. 2007](#); [Deegan et al. 2010](#)), the Colli Albani, Italy ([Freda et al. 2011](#)), Popocatépetl, Mexico ([Goff et al. 2001](#); [Schaaf et al. 2005](#)), and maybe even Yellowstone, USA ([Werner and Brantley 2003](#)), that like Vesuvius

are emplaced within some type of carbonate crust. We conclude that magma-carbonate interaction in the upper crust is likely a significant process operating beneath Vesuvius, but is variable in intensity. This interaction likely involves limestone and dolostone compositions and is not always easily quantified by conventional petrochemical indices (e.g., Sr-Ca relationships). Independently of the exact nature of the carbonate considered, our experiments demonstrate that magma-carbonate interaction can rapidly release considerable quantities of crustal CO₂ with potentially direct consequences on explosive versus effusive eruptive behaviour of the volcanic system.

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Figure Captions

Figure 1. BSE images of experimental products with increasing experimental durations ($t_d = 0$ to 300 s): limestone-bearing experiments (**a**, **b**, **c** and **d**) and dolostone-bearing experiments in (**e**, **f**, **g**, and **h**). Experiments show: *i*) a solid phase, i.e., calcite crystallite domains – marked with a broken white line in images **a**, **b** and **c** - and olivine and pyroxene crystals in images **f**, **g** and **h**; *ii*) a melt phase, i.e., Ca-normal, and Ca-rich and Ca-Mg-rich glasses; and *iii*) a vapour phase, preserved as bubbles. Note the contrasting brightness between Ca-rich and Ca-Mg-rich, and Ca-normal glasses in the images (light and dark colour, respectively).

Figure 2. BSE images showing dendritic micro-crystals of calcite in limestone-bearing experiments at $t_d = 60$ and 90 s (**a**, **b**), and clinopyroxene and olivine crystals from the dolostone-bearing experiments at $t_d = 90$ and 300 s (**c**, **d**). Abbreviations: Ol: olivine and Cpx: clinopyroxene.

Figure 3. Ternary diagrams of mineral compositions in dolostone-bearing experiments. Composition of pyroxenes in the $\text{CaSiO}_3 - \text{MgSiO}_3 - \text{FeSiO}_3$ system (**a**). Composition of olivines in the $\text{Ca}_2\text{SiO}_4 - \text{Mg}_2\text{SiO}_4 - \text{Fe}_2\text{SiO}_4$ system (small inset triangle). The bottom left hand corner of the small triangle is shown in large (**b**).

Figure 4. Representative microprobe chemical traverses in 0, 60, 90 and 300 s limestone-bearing experiments (**a** to **d**). The white lines in the images show the EMPA traverses (A to A') that correspond to the plots on the right. The Ca-normal and Ca-rich glasses are dark and pale grey in colour, respectively, and are separated by a dashed white line. The contamination front (CF) is the interfacial region between Ca-normal and Ca-rich domains. This area corresponds to a physico-chemical mixing and diffusion interface between the glass domains. Note that the width of the CF (delimited by vertical dashed lines on the right graphs) differs in the various experimental runs.

Figure 5. Representative microprobe chemical traverses in 0, 60, 90 and 300 s dolostone-bearing experiments (**a** to **d**). The solid white lines drawn on the images show the EMPA traverses (A to A') corresponding to the plots on the right. The contact between contaminated and uncontaminated melts is highlighted with white dashed lines (CF). See also **Fig 4**.

Figure 6. $^{87}\text{Sr}/^{86}\text{Sr}$ and CaO wt % profiles in the 60 s (**a**) and the 90 s (**b**) limestone-bearing experiments. The solid lines in the BSE images show different traverses in a single experiment (from A to A'). Drilled areas are numbered and highlighted in white on the BSE images. The right side of images for **a**) and **b**) correspond to the CaO wt. % from EMPA traverse (upper diagram) and the $^{87}\text{Sr}/^{86}\text{Sr}$ isotope ratios of the drilled areas (lower diagram). The height of the rectangles in the $^{87}\text{Sr}/^{86}\text{Sr}$ diagram represents the $\pm 2\text{SE}$ analytical uncertainty and their locations relative to the EMPA traverse data are approximate (see **Table 4**). The dark and pale shaded areas represent the glass domains observed in our experiments and the blue and green horizontal dashed lines are the CaO wt. % and $^{87}\text{Sr}/^{86}\text{Sr}$ ratios of the starting materials.

Figure 7. $^{87}\text{Sr}/^{86}\text{Sr}$ and CaO wt. % profiles in the dolostone-bearing experiments. **a**) BSE image of the 60 s experiment and **b**), **c**) and **d**) show the 90 s experiment. The white lines in the images show different traverses in a single experiment (A to A'). The diagrams on the right side illustrate CaO wt. % (upper diagram) and $^{87}\text{Sr}/^{86}\text{Sr}$ (lower diagram) for the different traverses and drill holes, respectively. The height of the rectangles in the $^{87}\text{Sr}/^{86}\text{Sr}$ diagram includes the $\pm 2\text{SE}$ analytical uncertainty and their locations of the drill holes relative to the EMPA traverses are approximate. The traverse in panel **b**) extends beyond the BSE image. Abbreviations as in **Fig 6**.

Figure 8. Compositional summary diagram of all limestone-bearing experiments. Binary mixing trajectory between shoshonite and limestone is illustrated. The experiments are represented by: triangles for $t_d = 0$ s ($n = 17$); diamonds for $t_d = 60$ s ($n = 27$); squares for $t_d =$

90 s ($n = 21$); circles for $t_d = 300$ s ($n = 11$). The glass domains and the interface region are given as: dark grey = Ca-rich, pale grey = Ca-normal, and white = contamination front. The maximum degree of chemical mixing of ambient melt with carbonate derived CaO is reached at $t_d = 90$ s and equates to about 60 %. Note that at $t_d = 300$ s (highlighted with a red line), the system has progressively homogenised (less than 25 % carbonate derived CaO in mixture). The small displacement of CaO and SiO₂ above the linear binary mixing trajectory reflects fluctuations of these elements as a consequence of transport through convection and interdiffusion between the experimental melts (see text for details and **Online Resource 1 Fig OR1**).

Figure 9. Compositional summary diagram of all dolostone-bearing experiments. Binary mixing trajectory between shoshonite and dolostone is represented. The symbols for each experiment and the colours of glass domains are the same as in **Fig 8**. The total number of data points in each experiment is: $n = 12$ for $t_d = 0$ s; $n = 80$ for $t_d = 60$ s; $n = 137$ for $t_d = 90$ s; $n = 23$ for $t_d = 300$ s. For these experiments, the system reached its maximum degree of mixing at $t_d = 60$ to 90 s. For $t_d = 300$ s (highlighted with a red line) the glass has again adopted a more homogenous composition. The deviations from the binary mixing trajectory is likely a coupled consequence of melt transport, as in the limestone-bearing experiments, but was affected by simultaneous crystal growth and associated Mg removal from the melt (see also **Online Resource 1 Fig OR2**).

Figure 10. a) $^{87}\text{Sr}/^{86}\text{Sr}$ values for our experimental glasses in comparison to VVS shoshonite magma compositions, Vesuvius skarn xenoliths, and local carbonate crust. The Ca-normal glass is within the range of the natural shoshonite magmas, whereas the contaminated glass is displaced towards crustal values and overlaps with the range of skarns reported from the VVS. **b)** $^{87}\text{Sr}/^{86}\text{Sr}$ vs. Sr (ppm) diagram illustrates possible binary mixing relationships. Triangles and circles represent the limestone-bearing experiments for t_d at 60 and 90 s, respectively, whereas squares and diamonds are the dolostone-bearing experimental products for the same

experimental durations. The mixing trajectories A-C₁ and A-C₂ define the mixing space between the shoshonite end-member (A) and the high ⁸⁷Sr/⁸⁶Sr and high Sr concentrations reported from some parts of the nearby carbonate crust. The mixing trajectories A-C₂, A-C₃, A-C₄ and A-C₅ form the mixing space between shoshonite (A) and the low ⁸⁷Sr/⁸⁶Sr and low-Sr ppm carbonates reported from the Campanian region (see text for details). The majority of the drilled experimental glass compositions fall within the low Sr-carbonate mixing space (mixing curves A-C₂ to A-C₅), implying mixtures of between 55 and 75% carbonate-derived ⁸⁷Sr/⁸⁶Sr relative to shoshonite. Literature data from: [Civetta et al. \(1991\)](#); [Di Renzo et al. \(2007\)](#); [Del Moro et al. \(2009\)](#); [Piochi et al. \(2006\)](#); [Iannace et al. \(2011\)](#).

Table 1: Starting material composition

	Hydrous glass ^(a)	1 σ (9) ^(b)	Limestone CaVe1 Procida ^(c)	Dolostone CaVe3 Procida ^(d)
SiO ₂	49.85	0.21	0.02	0.02
TiO ₂	1.03	0.08	-	-
Al ₂ O ₃	15.95	0.67	-	-
FeO _t ^(e)	7.98	0.25	0.01	0.0
MnO	0.14	0.02	0.01	0.01
MgO	6.03	0.33	0.79	21.01
CaO	9.98	0.29	54.99	31.34
Na ₂ O	2.35	0.07	-	-
K ₂ O	4.01	0.19	-	-
P ₂ O ₅	0.70	0.09	0.02	0.0
CO ₂	-	-	44.11*	47.59*
H ₂ O	2		-	-
Total	100.02		100.00	100.00
Sr	735 [†]		299 ^{††}	89 ^{††}
⁸⁷ Sr/ ⁸⁶ Sr ^{†††}	0.706661 ±8		0.7075 ±1	0.7072 ±1

^(a) Hydrous glass synthesised from natural powdered rock at T = 1250 °C and 0.5 GPa.

^(b) 1 σ standard deviation; the number in parenthesis represents the number of analyses.

^(c) Lower Cretaceous limestone from Procida Formation.

^(d) Triassic dolostone from Procida Formation.

^(a, c, d) Major elements composition of the starting material were determined by electron-microprobe at INGV and given in wt. %.

^(e) Total iron is given as FeO_t.

*Fraction of CO₂ in the fluid phase was calculated assuming wt. % CaCO₃ in the samples as equivalent to 100 wt. % minus all other element oxides.

[†] Strontium content for shoshonitic melt: Di Renzo et al. (2007).

^{††} Limestone and dolostone Sr contents were determined by ICP-emission spectrometry at Acmelabs (Vancouver, Canada) and are given in ppm.

^{†††} Strontium isotopes for: shoshonitic melt: Di Renzo et al. (2007); and for limestone and dolostone: Piochi et al. (2006) and Iannace et al. (2011).

Table 2: Experimental Conditions

Run #	Experimental time t_d (s)	Pressure (GPa)*	Temperature (°C)*	Limestone (mg)	Dolostone (mg)	Hydrous glass ~ 2 wt. % H ₂ O (mg)
PC443-V5	0	0.5	1200	6.1	-	29.6
PC448-V9	60	0.5	1200	8.2	-	33.3
PC442-V3	60	0.5	1200	6.3	-	28.8
PC441-V1	90	0.5	1200	6.8	-	24.2
PC-445 V7	300	0.5	1200	7.2	-	26.0
PC443-V6	0	0.5	1200	-	6.0	30.0
PC448-V10	60	0.5	1200	-	7.7	34.2
PC442-V4	60	0.5	1200	-	7.0	29.6
PC441-V2	90	0.5	1200	-	6.0	30.0
PC445-V8	300	0.5	1200	-	7.1	26.4

*Pressure and temperature uncertainties are ± 0.02 GPa and ± 5 °C, respectively.

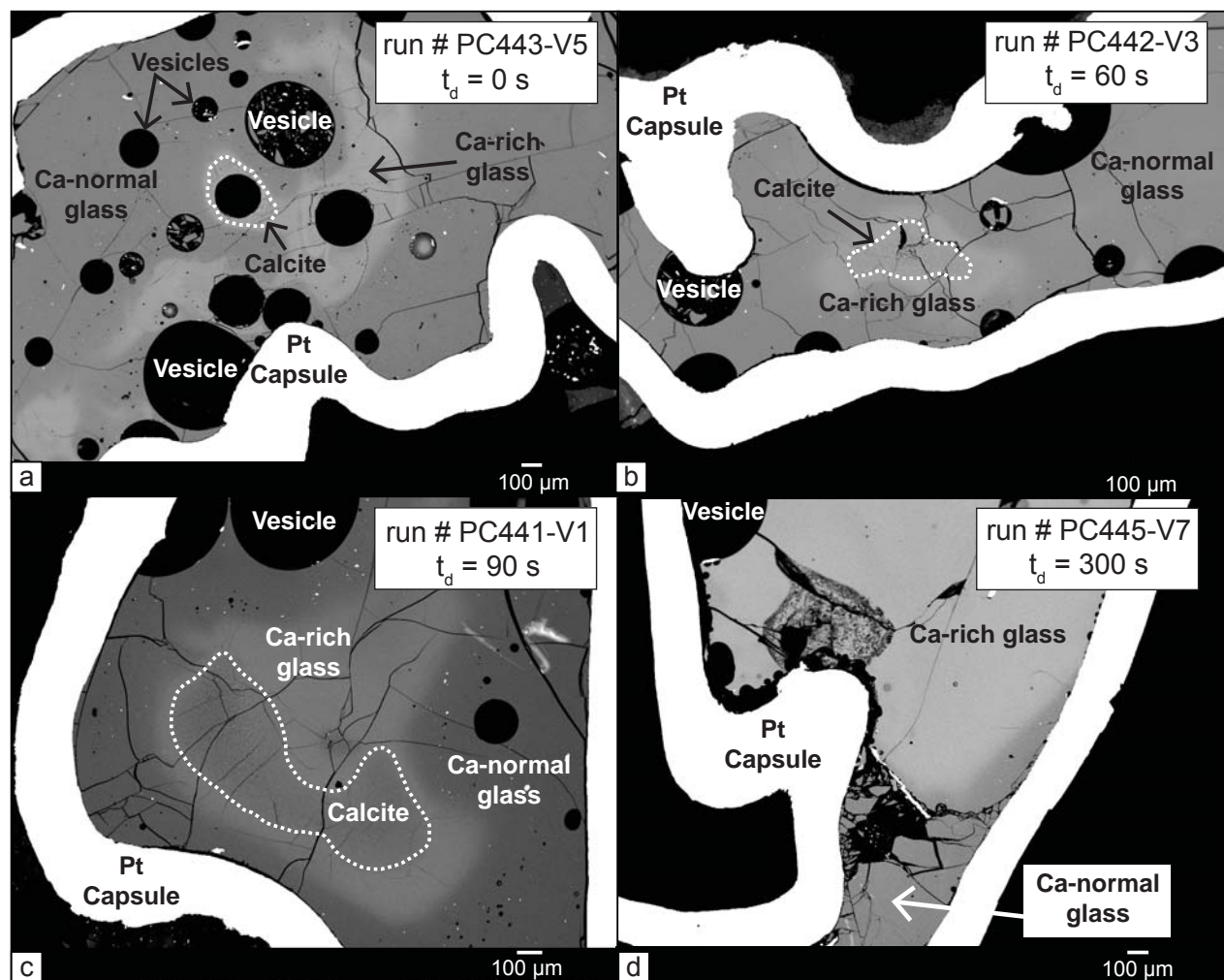
Table 3 : Continued

Experimental run # PC442-V4 Dolostone-bearing																						
Traverse 1											Traverse 2											
Ca-Mg-normal glass											Contamination front (CF)			Ca-Mg-rich glass			Contamination front (CF)					
SiO ₂	48.9	48.8	49.0	48.7	48.6	44.1	40.9	41.6	38.5	37.0	45.1	47.9	46.5	43.9	47.0	46.6						
Al ₂ O ₃	1.1	0.9	0.9	1.0	0.9	1.0	1.1	0.7	0.8	0.7	1.0	1.0	0.9	1.0	1.0	0.9						
TiO ₂	16.7	16.6	16.6	16.3	16.4	14.7	13.8	14.0	12.7	12.1	15.4	16.4	15.8	15.8	16.2	16.3						
FeO	6.5	6.3	6.6	6.2	6.1	6.6	6.9	6.1	5.7	5.4	5.7	5.3	5.5	5.7	5.5	5.5						
MnO	0.1	0.1	0.1	0.1	0.1	0.1	0.2	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1						
MgO	5.8	5.7	5.7	5.8	5.8	7.8	8.7	8.4	9.2	9.0	7.1	5.5	7.2	7.8	6.6	6.7						
CaO	9.1	9.2	9.1	9.4	10.2	14.4	16.6	17.1	19.9	20.8	13.4	10.1	12.6	14.9	10.5	11.2						
Na ₂ O	3.0	2.9	3.1	3.1	2.9	2.2	1.9	1.9	1.6	2.4	2.4	2.9	2.6	2.3	3.0	3.0						
K ₂ O	3.9	4.0	4.0	4.2	4.1	2.6	2.2	2.6	2.1	2.3	3.7	4.9	3.6	3.0	4.3	4.2						
P ₂ O ₅	0.6	0.5	0.6	0.6	0.7	0.6	0.6	0.6	0.6	0.5	0.5	0.7	0.5	0.6	0.6	0.6						
BaO	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.0	0.1	0.0	0.1	0.0	0.2	0.1	0.1						
SrO	0.2	0.1	0.2	0.0	0.1	0.1	0.2	0.2	0.0	0.1	0.3	0.3	0.2	0.1	0.1	0.1						
Total*	96.1	95.3	96.0	95.4	95.9	94.2	93.1	93.4	91.3	90.4	94.8	95.2	95.7	95.5	95.0	95.5						

Note: analyses shown are from the experimental glass regions: Ca-normal, Ca-rich and contamination front (CF).

*The low totals obtained are due to one of the following reasons: *a*) microbubbles (cf. Deegan et al., 2010), *b*) increased CO₂ solubility in alkaline and ultrapotassic melts in general (cf. Behrens et al., 2009) or *c*) increased CO₂ solubility in our Ca-rich experimental melt (Moore, 2008).

Limestone-bearing



Dolostone-bearing

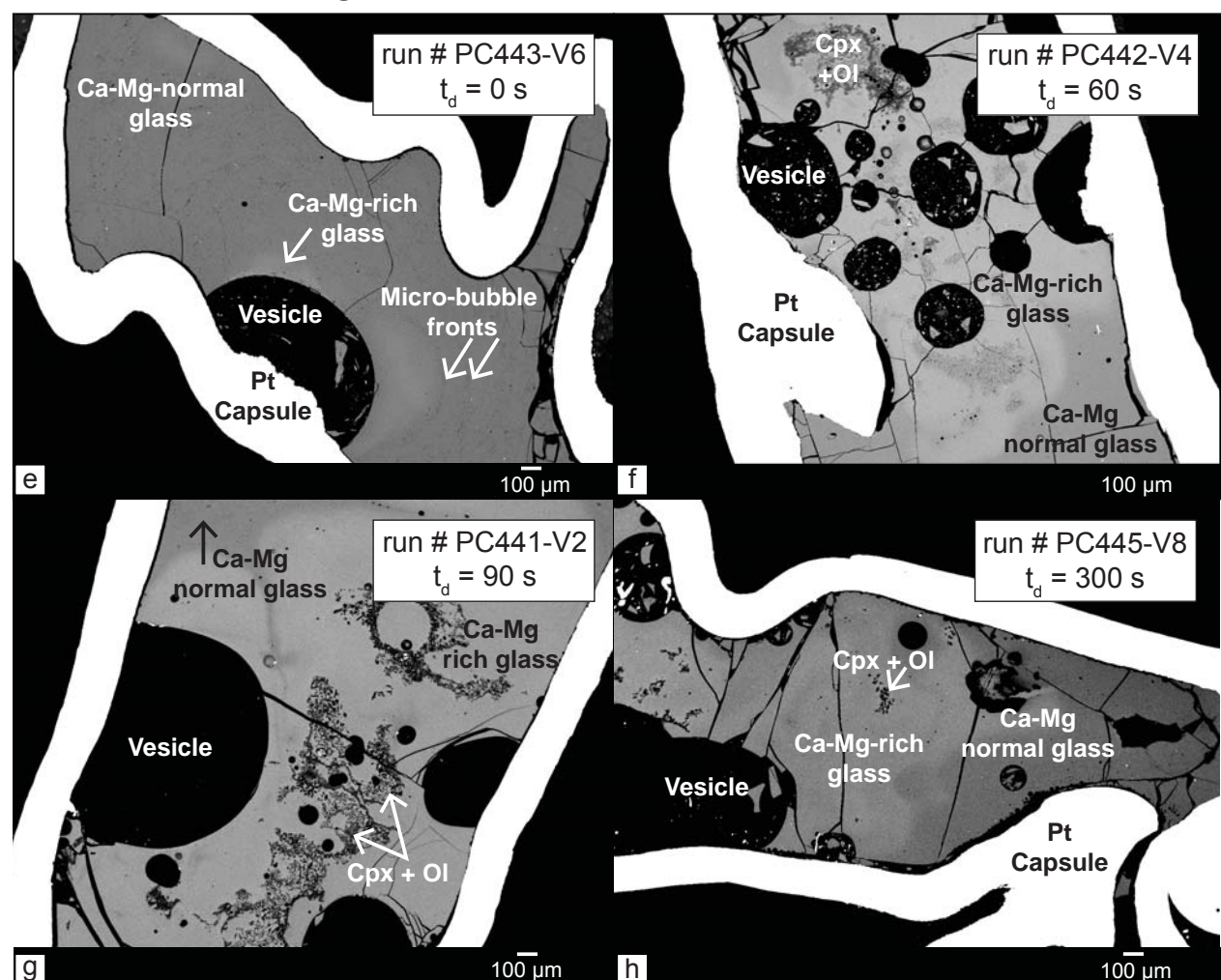


Figure 1

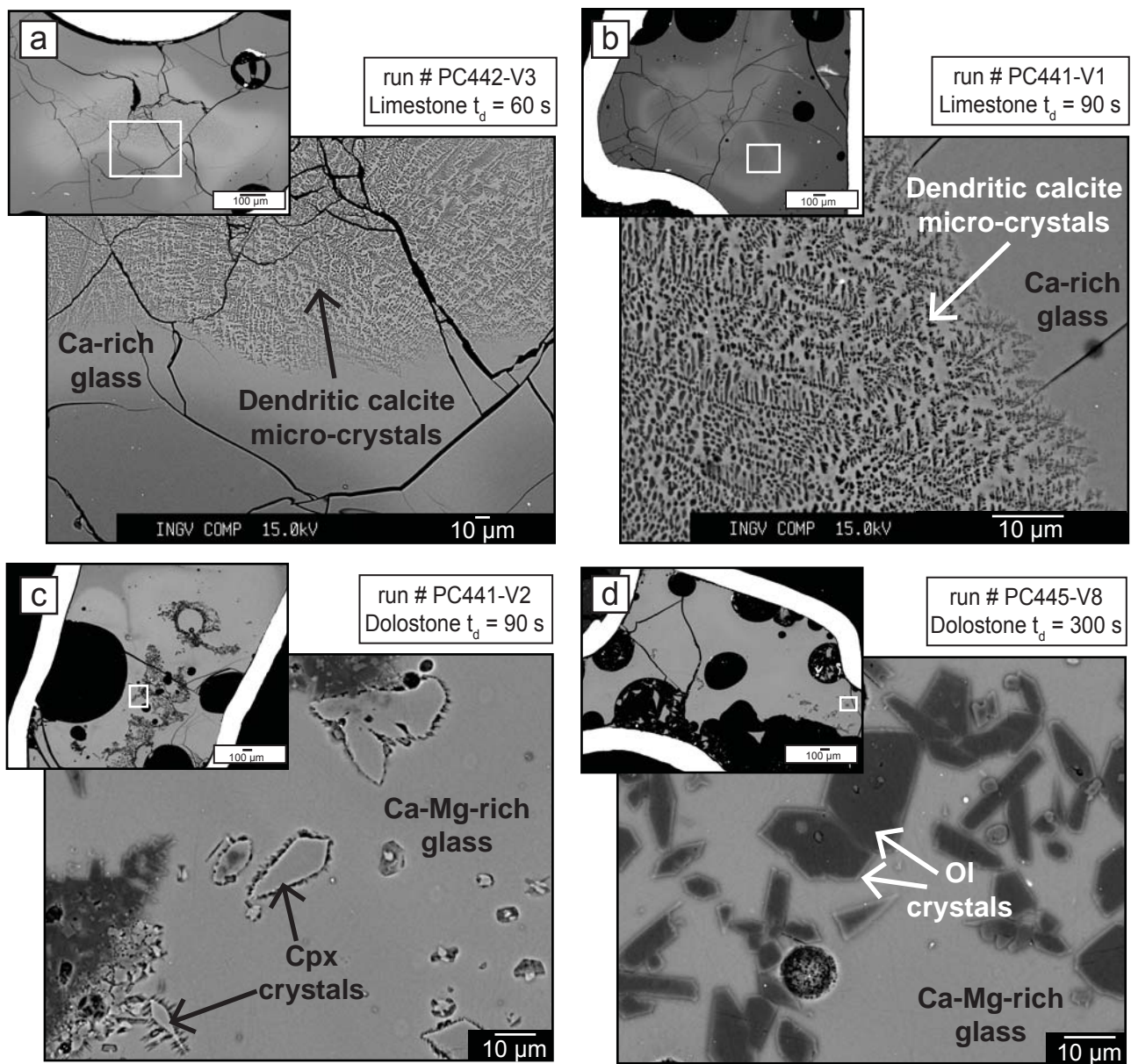


Figure 2

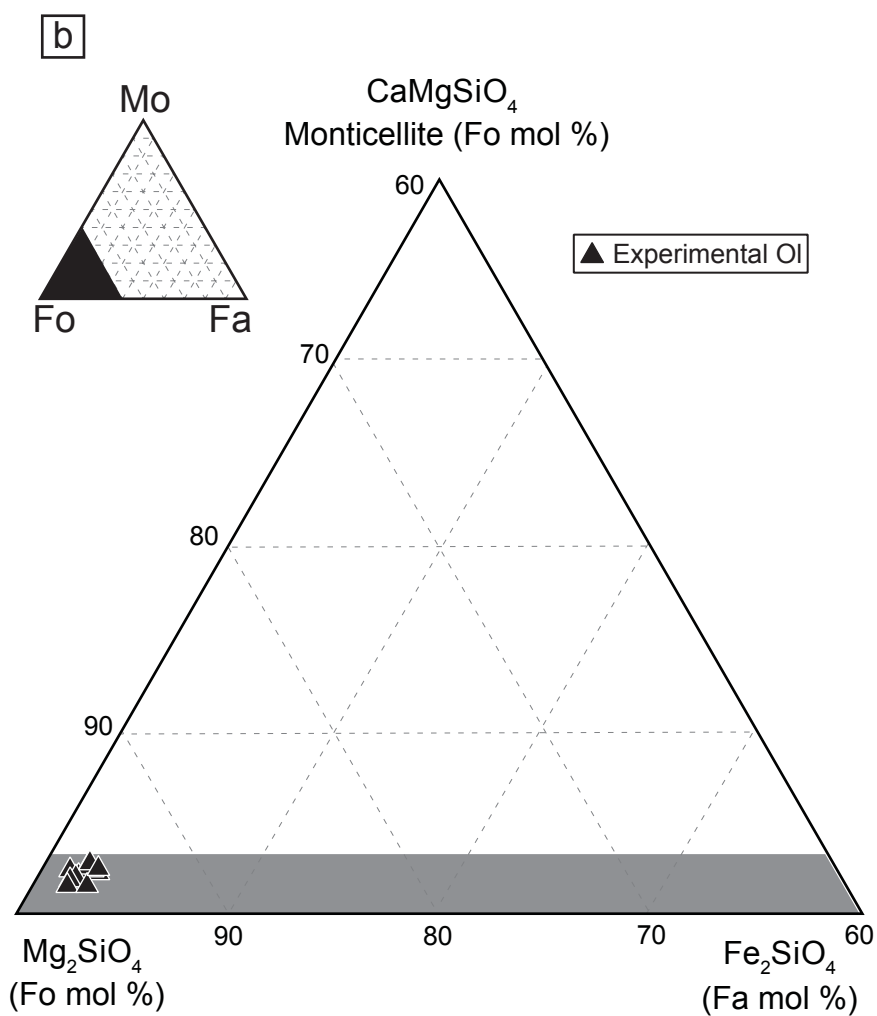
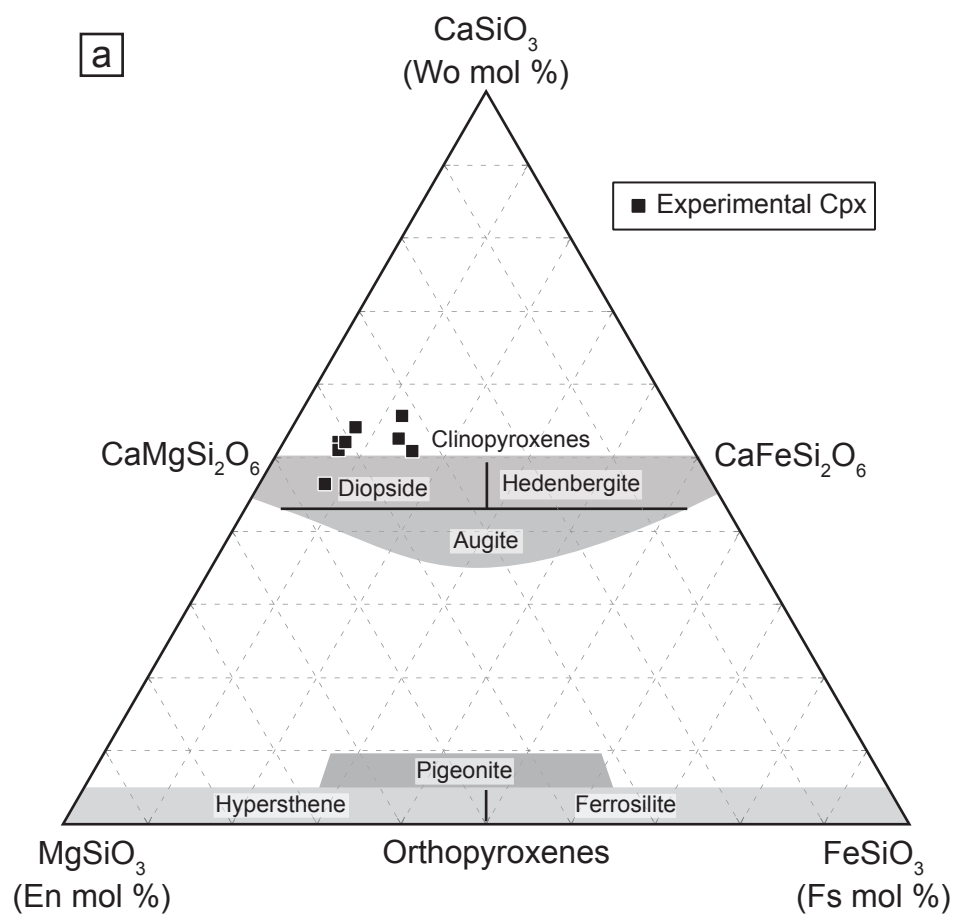
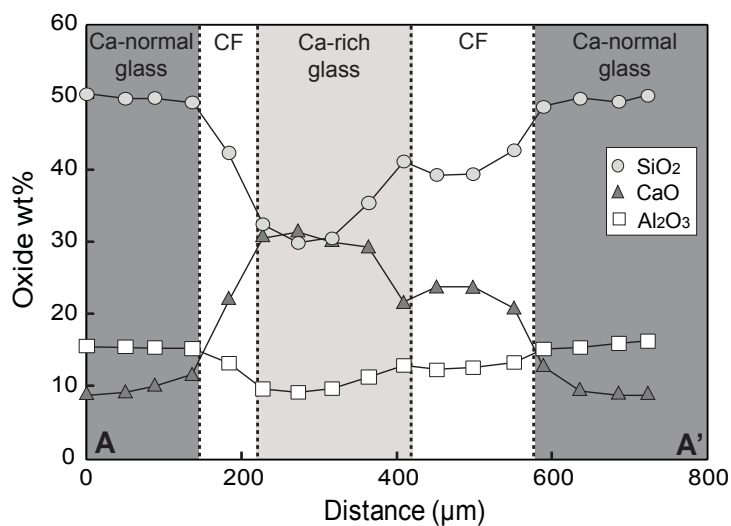
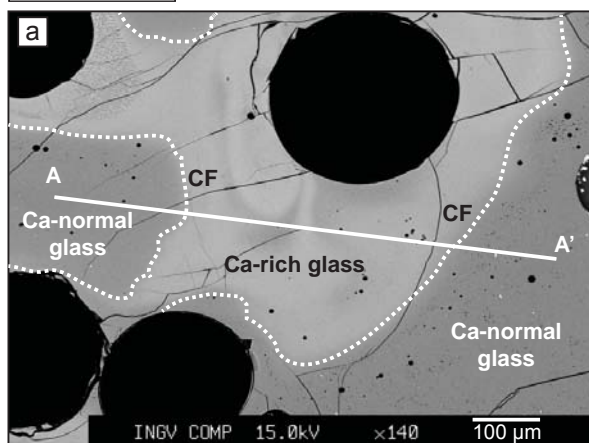
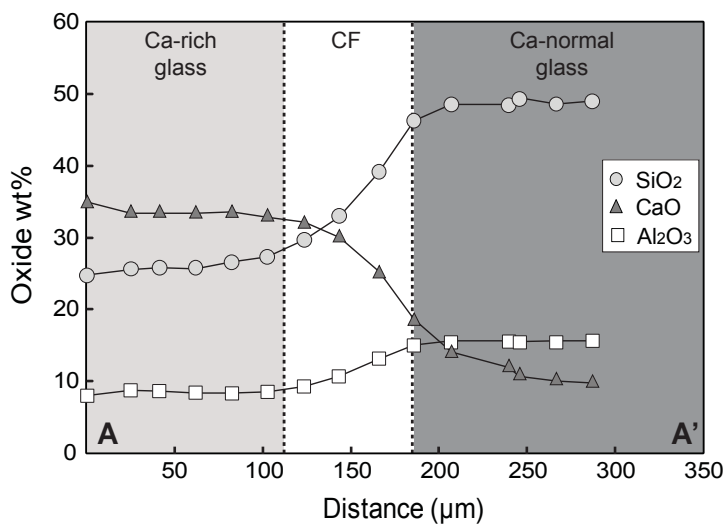
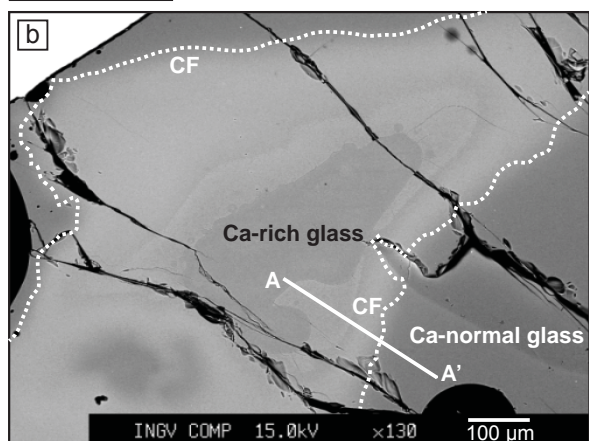


Figure 3

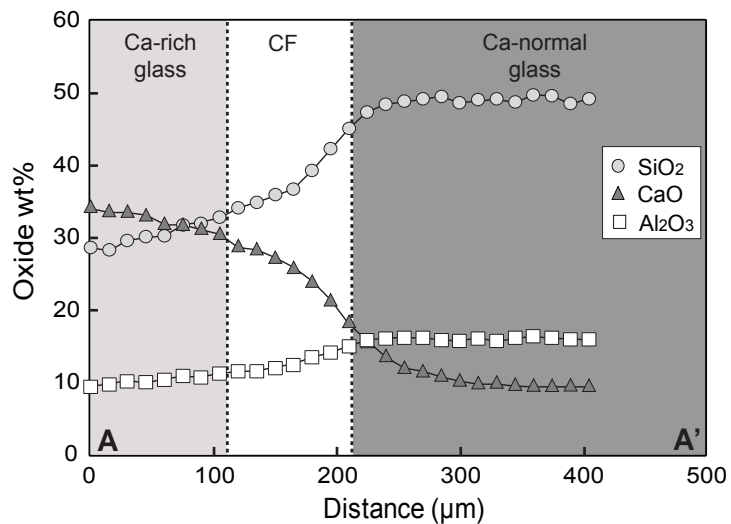
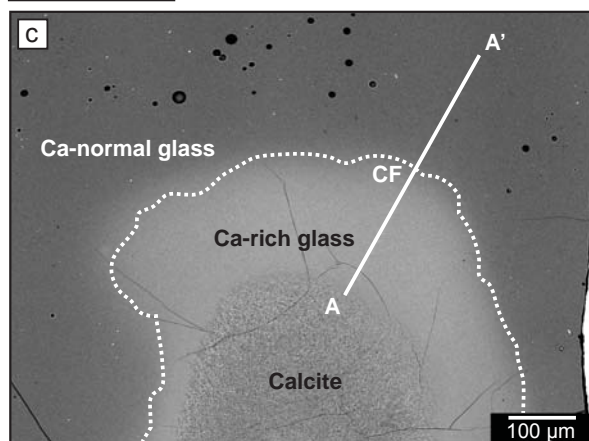
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run # PC448-V9
t_d = 60 s



run # PC441-V1
t_d = 90 s



run # PC445-V7
t_d = 300 s

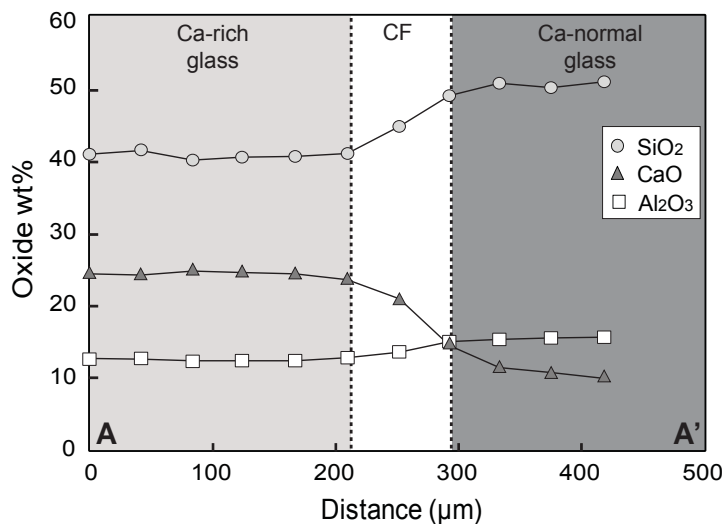
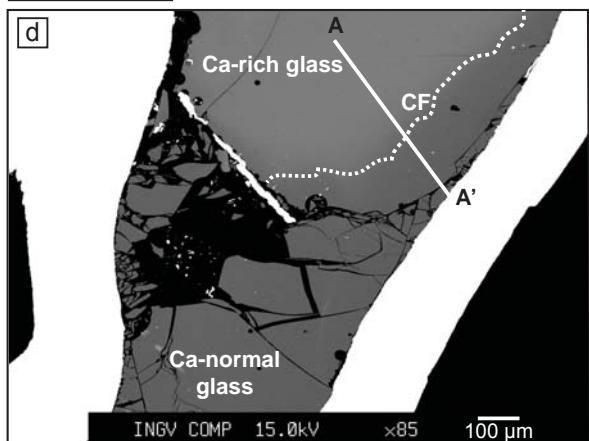


Figure 4

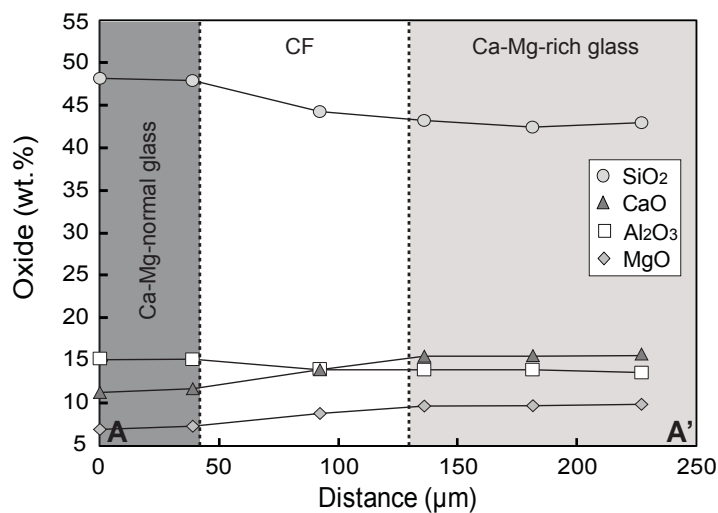
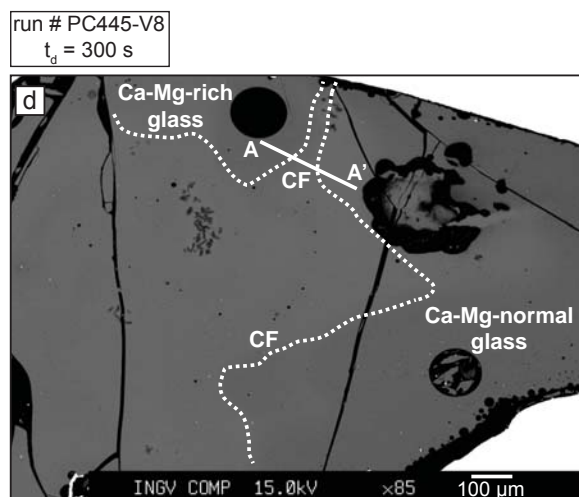
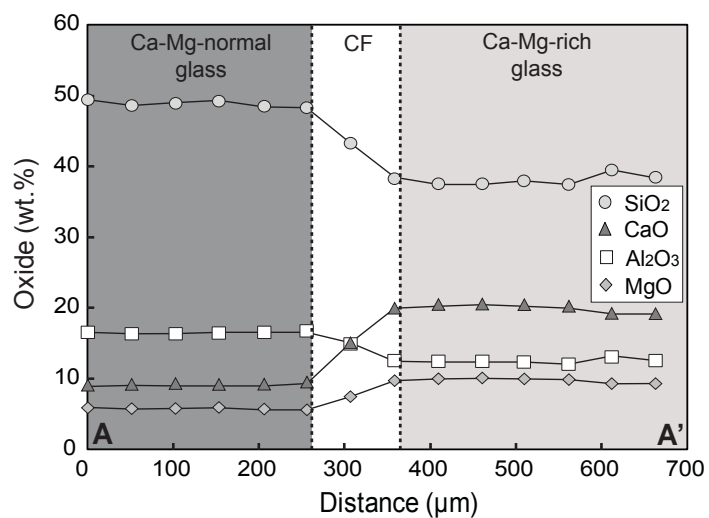
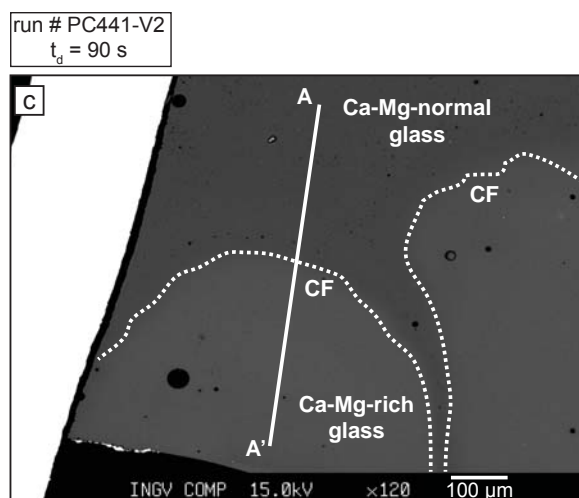
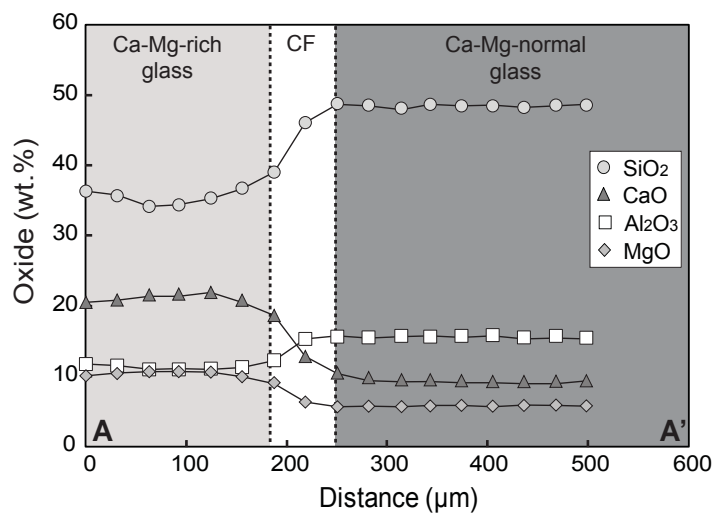
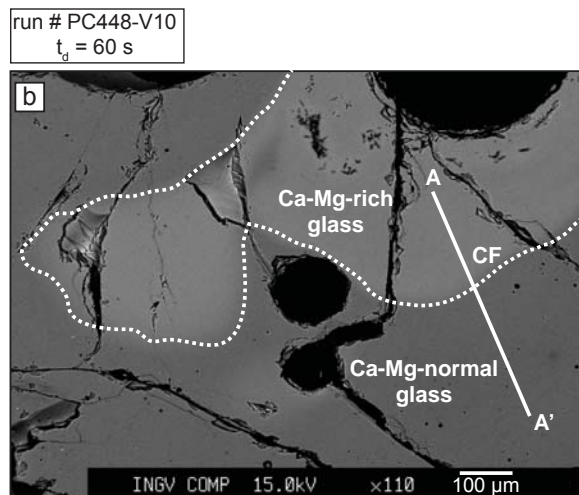
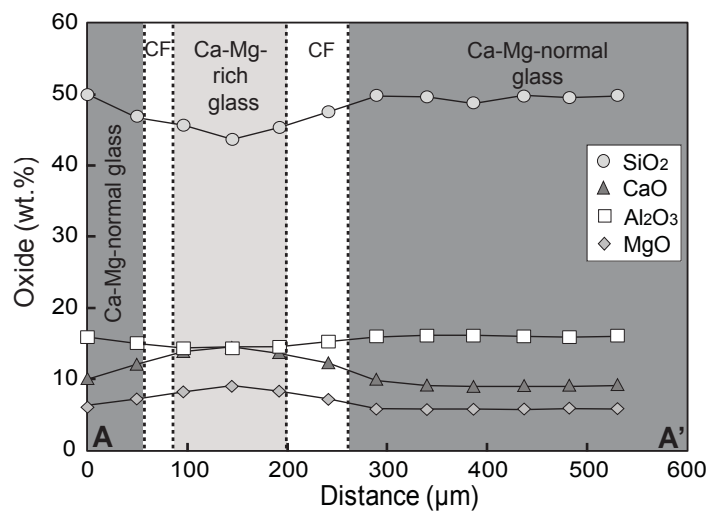
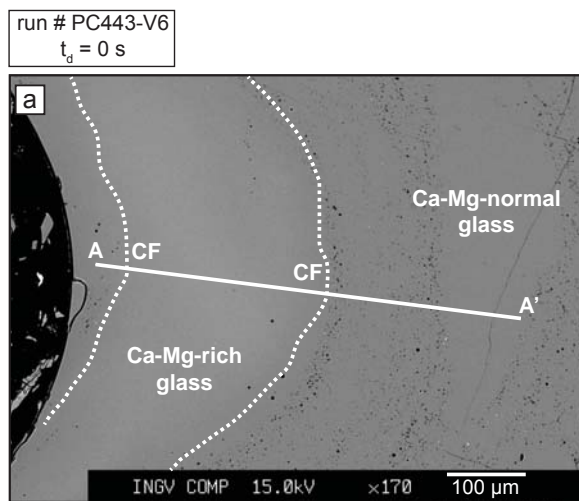


Figure 5

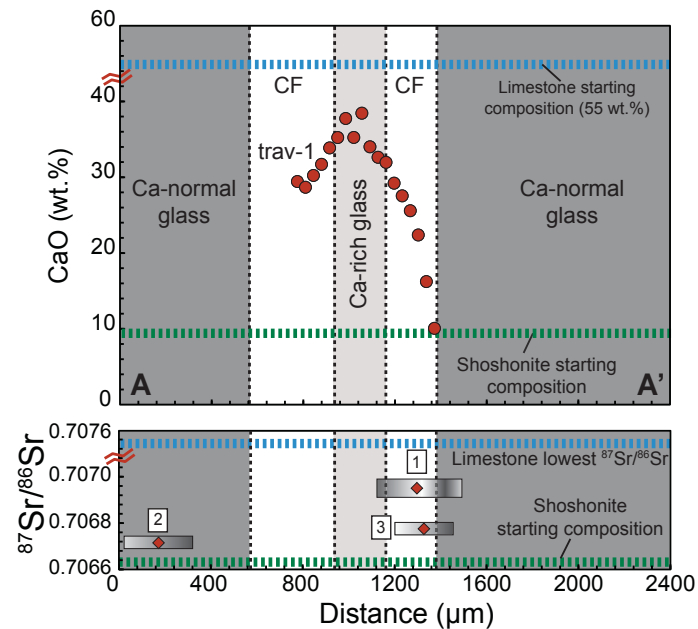
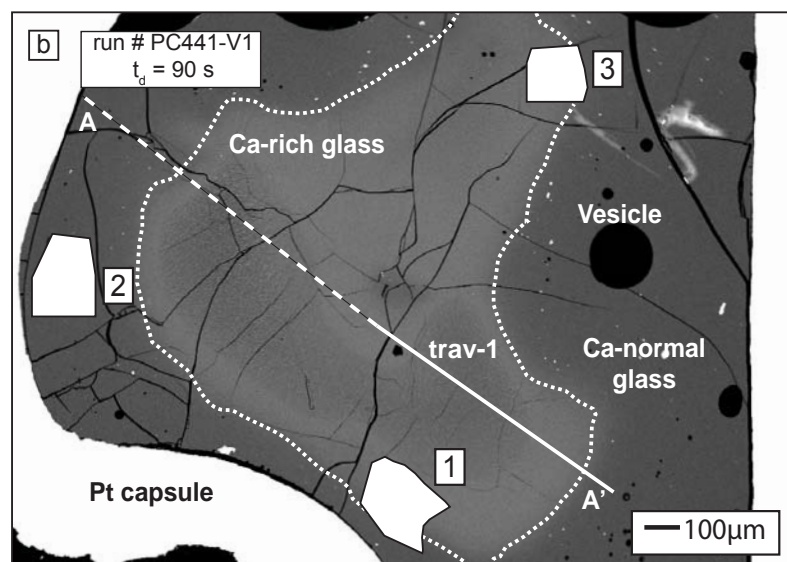
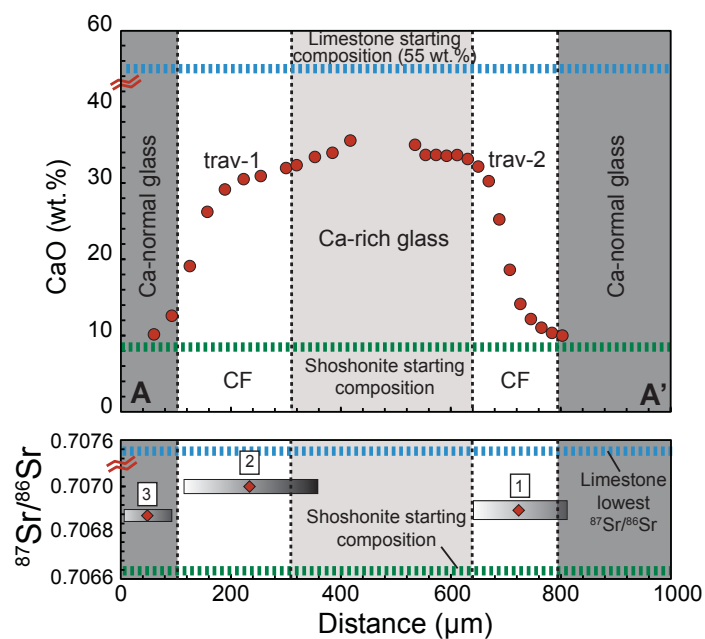
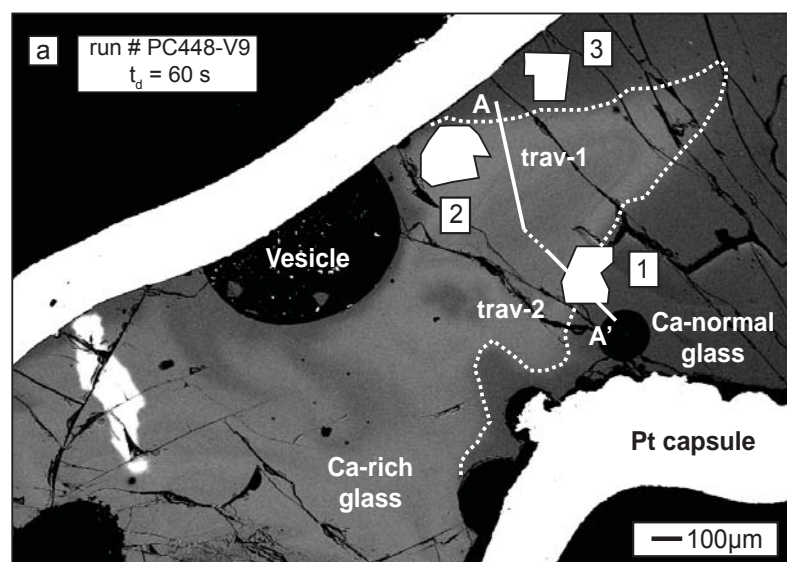


Figure 6

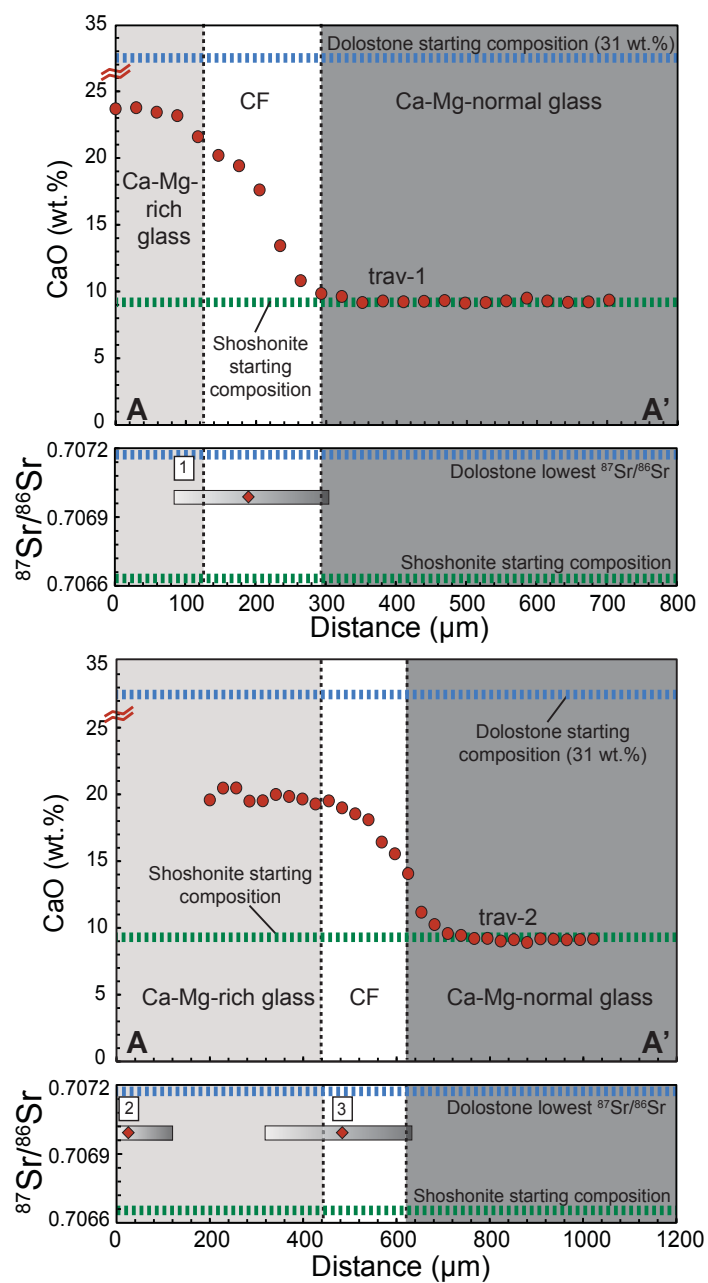
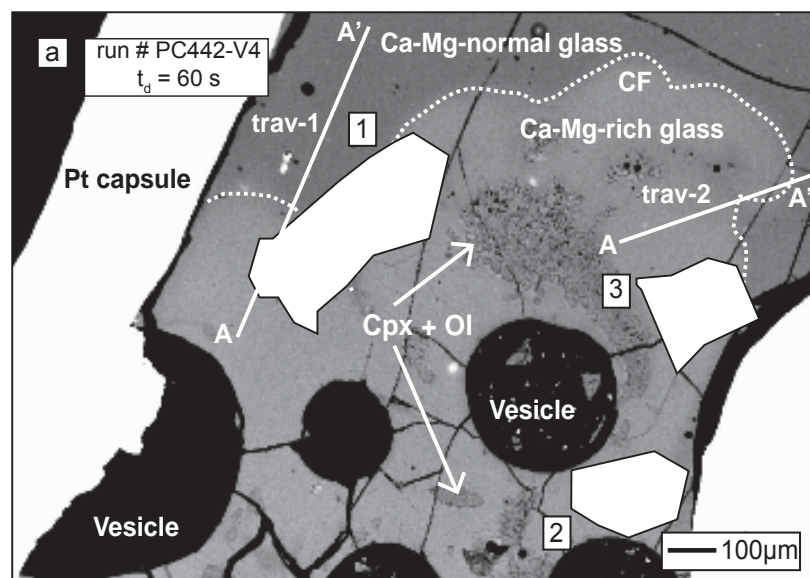


Figure 7

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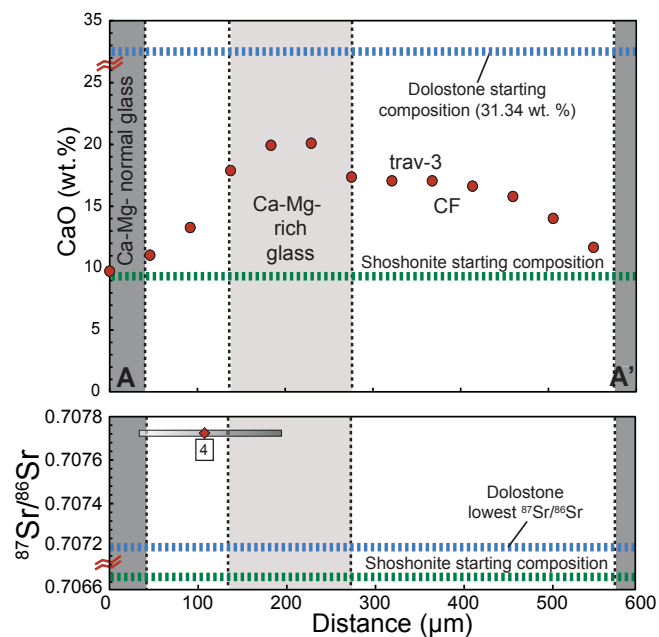
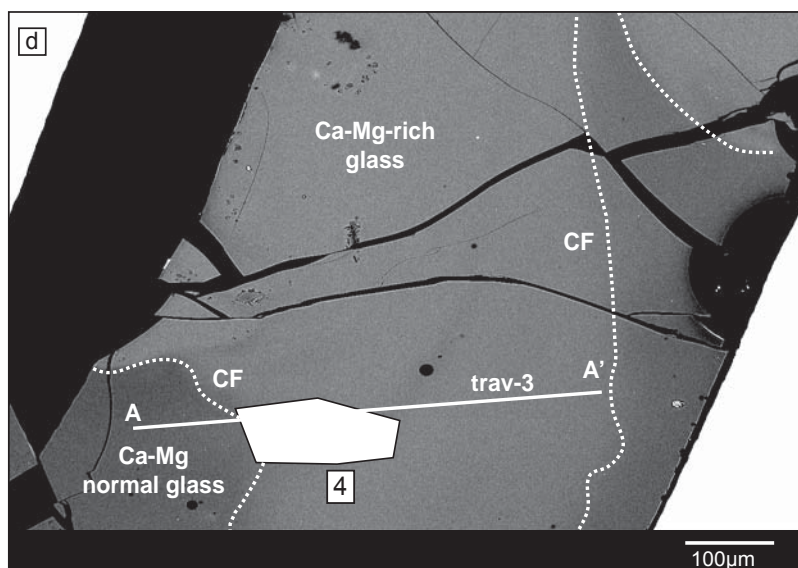
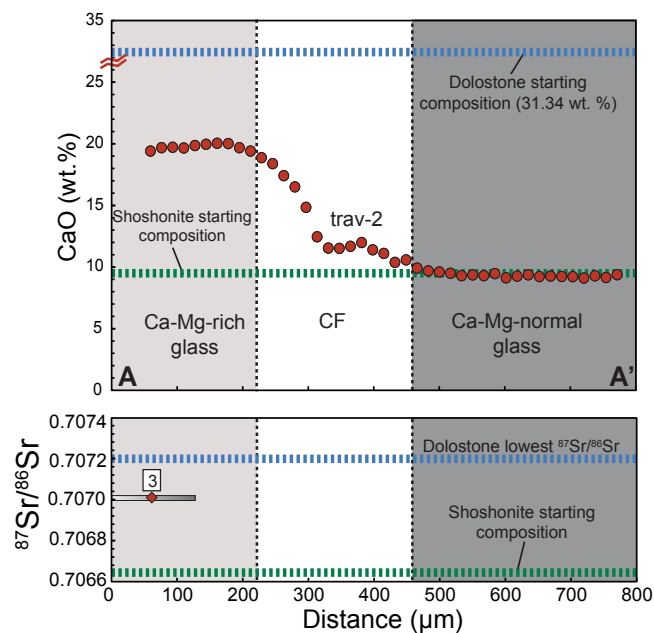
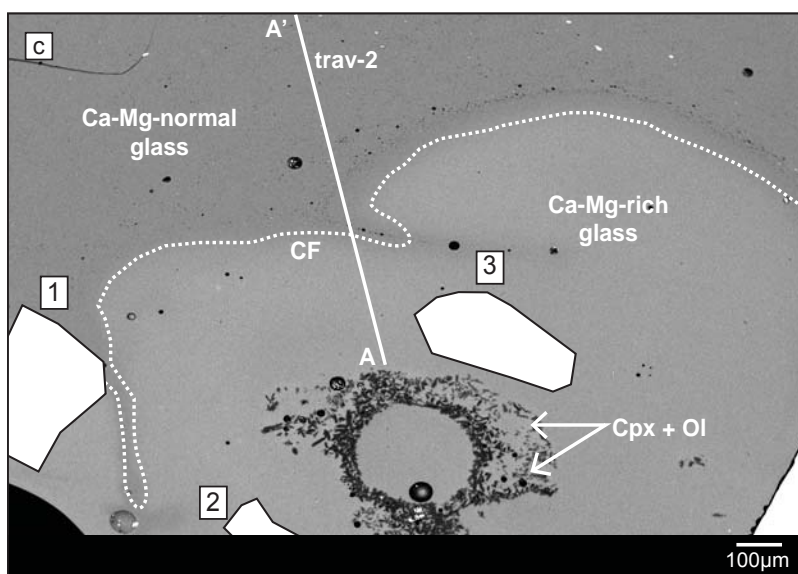
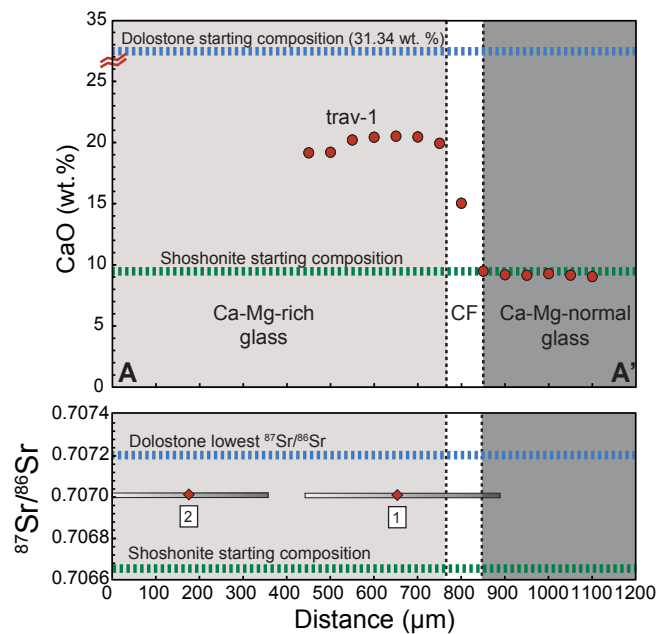
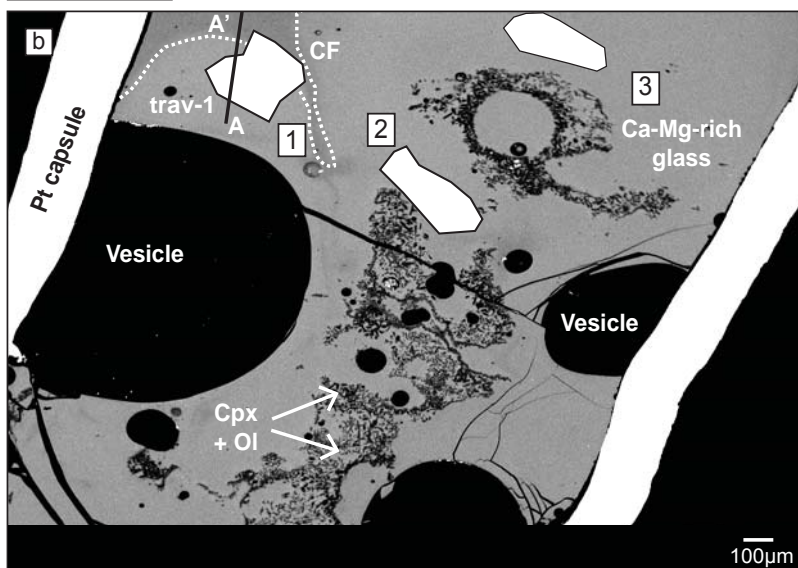


Figure 7 continued

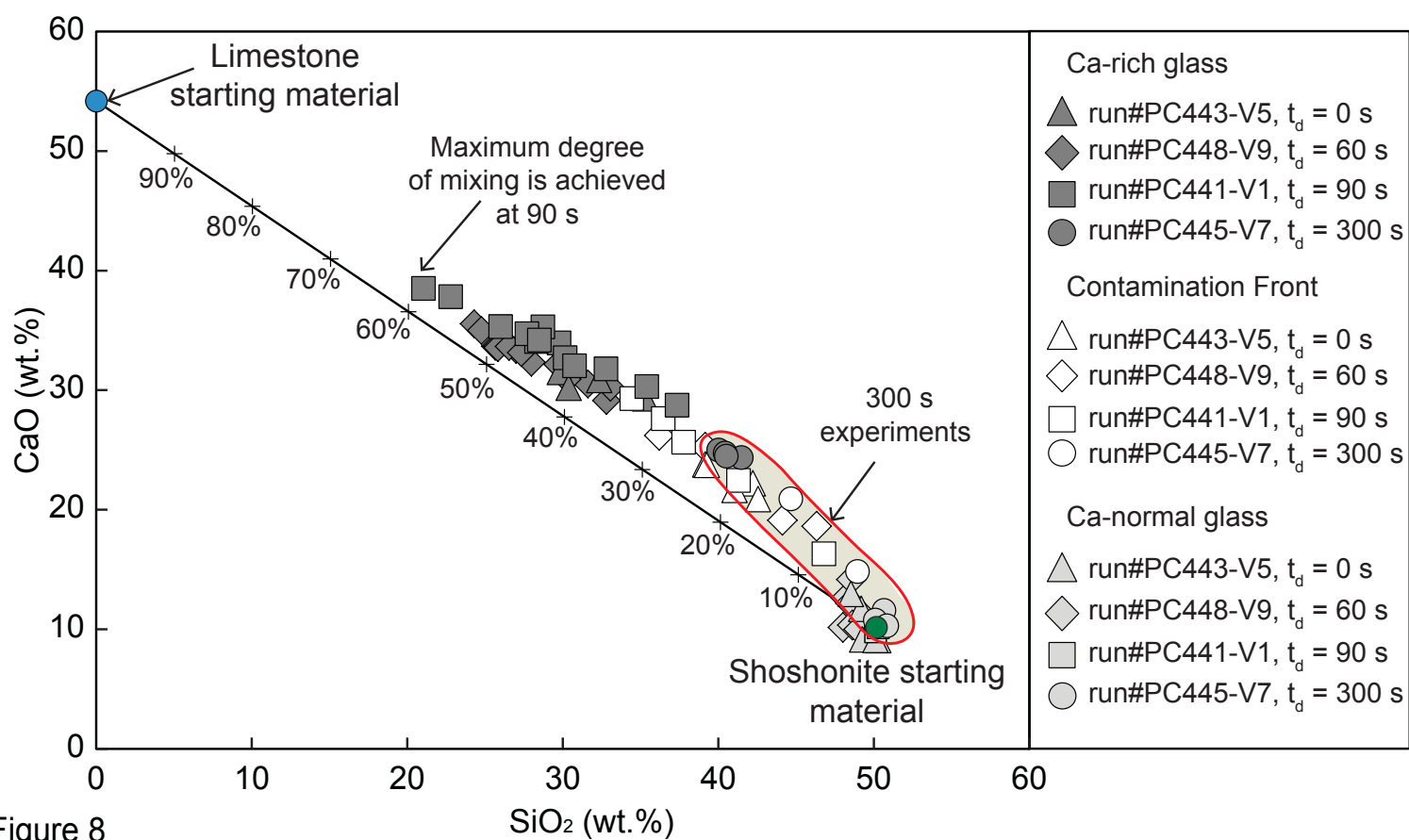


Figure 8

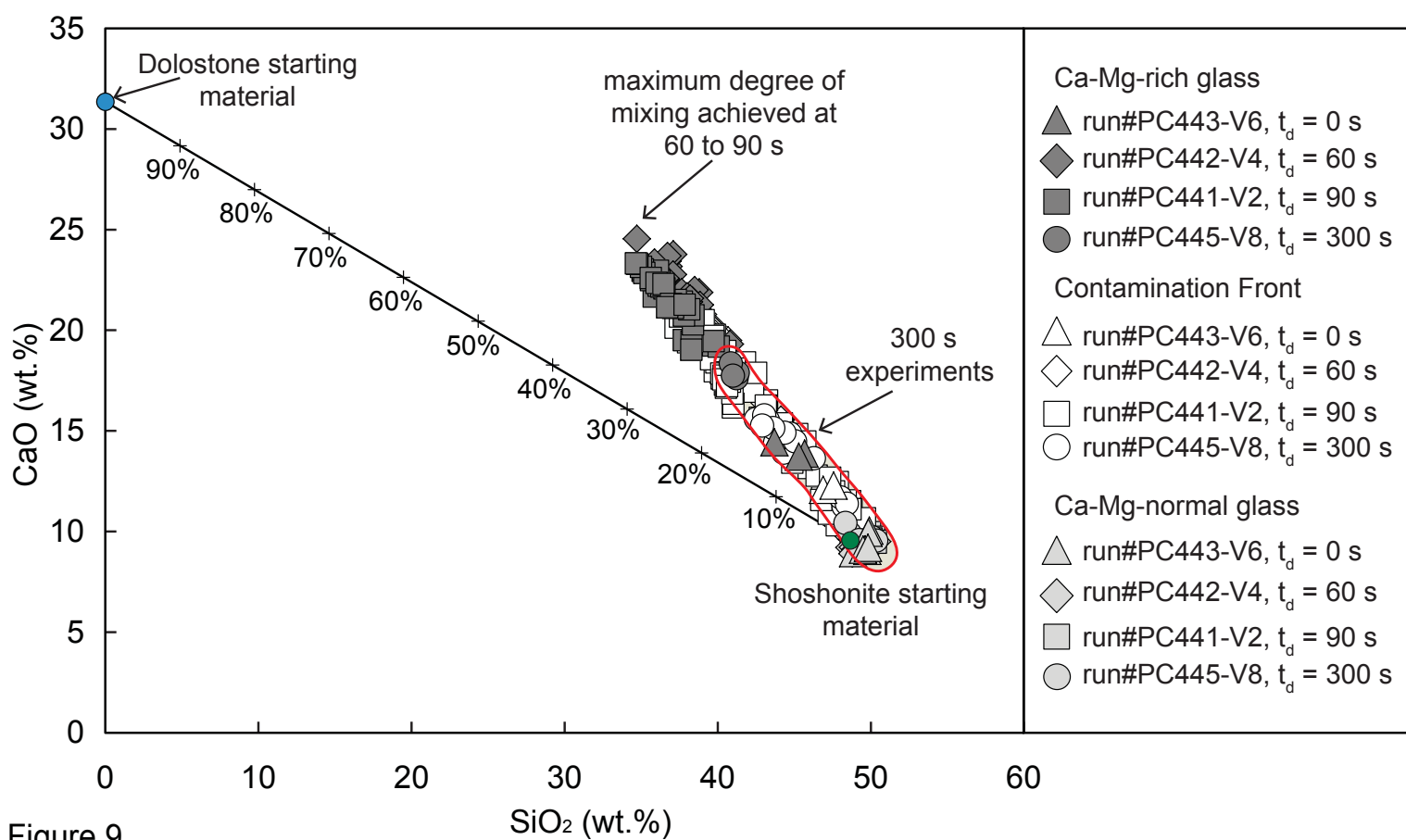


Figure 9

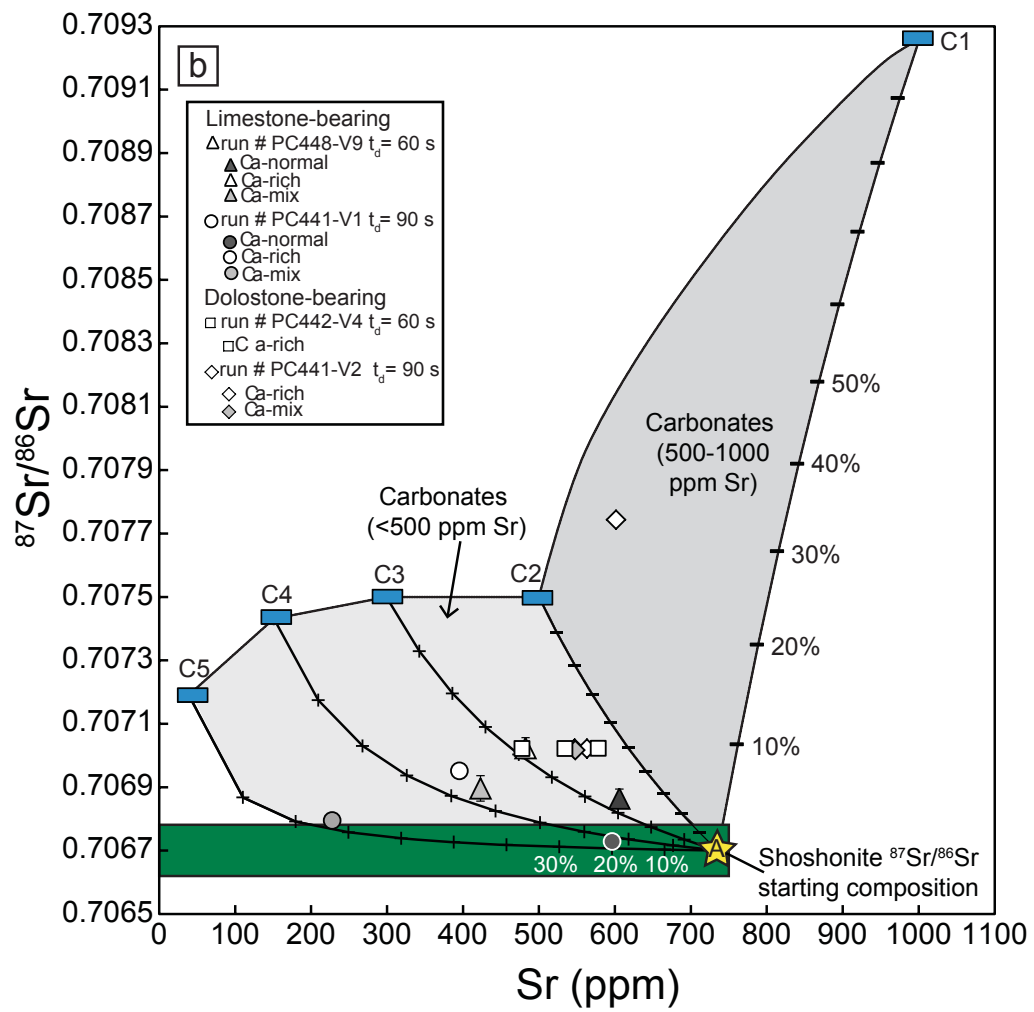
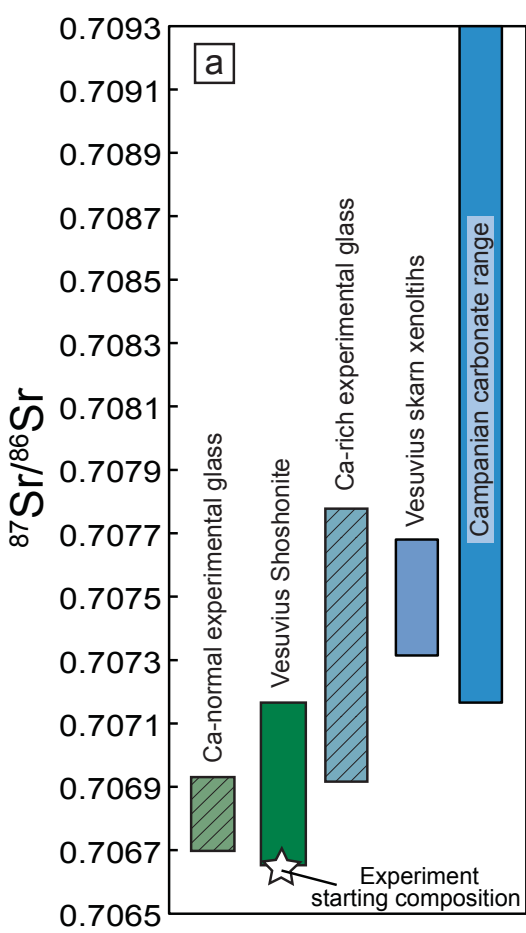


Figure 10

Online Resource Table OR1

Electron Microprobe chemical analysis for pyroxene and olivine.
Concentrations are given in mol % as shown in **Fig 3**.

Table 1

Experiment	PC441-V4 (60 s)								
	Px 1	Px 2	Px 3	Px 4	Px 5	Px 6	Px 7	Px 8	Px 9
SiO ₂	39.1	39.9	37.3	44.2	44.7	40.2	43.1	44.4	44.8
TiO ₂	2.0	2.1	2.1	1.4	1.3	1.3	1.6	1.4	1.3
Al ₂ O ₃	16.4	16.2	17.0	13.1	13.0	17.6	14.2	13.2	12.9
FeO	8.3	7.7	9.8	4.8	4.7	5.0	4.8	4.4	4.5
MnO	0.1	0.0	0.0	0.0	0.1	0.1	0.0	0.1	0.0
MgO	10.7	10.0	10.4	12.7	13.2	14.4	11.8	13.0	13.2
CaO	23.7	25.0	23.0	23.3	23.0	20.7	23.8	23.3	23.3
Na ₂ O	0.2	0.2	0.2	0.2	0.2	0.1	0.1	0.2	0.1
P ₂ O ₅	0.6	0.7	0.7	0.4	0.3	0.4	0.3	0.4	0.3
Total	101.2	101.8	100.4	100.2	100.4	99.9	99.8	100.3	100.4
Wo	52.60	55.66	50.89	52.19	51.10	46.47	54.19	52.10	51.73
En	33.00	30.97	32.21	39.43	40.74	44.77	37.21	40.31	40.57
Fs	14.41	13.38	16.89	8.38	8.16	8.75	8.60	7.59	7.70

Table 1 (continued)

Experiment	PC441-V4 (60 s)		PC441-V2 (90 s)				PC445-V8 (300 s)				
	Ol 1	Ol 2	Ol 3	Ol 4	Ol 5	Ol 6	Ol 7	Ol 8	Ol 9	Ol 10	Ol 11
SiO ₂	41.0	40.8	40.8	41.7	41.4	41.7	41.1	41.2	41.1	40.2	41.7
TiO ₂	0.1	0.2	0.3	0.1	0.2	0.1	0.1	0.1	0.2	0.3	0.1
Al ₂ O ₃	0.8	1.8	0.9	0.6	0.5	0.4	0.7	0.4	0.7	1.1	0.5
FeO	1.7	1.6	1.6	2.6	2.0	2.6	2.2	2.0	1.3	2.5	2.8
MnO	0.1	0.0	0.1	0.1	0.1	0.0	0.1	0.0	0.1	0.0	0.1
MgO	55.2	55.5	55.3	53.9	54.4	55.1	54.7	54.8	54.9	54.3	53.7
CaO	1.8	1.6	1.3	2.1	2.4	1.4	2.0	1.6	1.8	1.9	1.7
Na ₂ O	0	0	0	0.02	0	0.003	0.012	0	0	0.002	0
P ₂ O ₅	0.07	0.06	0.07	0.02	0.04	0.06	0.05	0	0.02	0.04	0
Total	100.7	101.5	100.3	101.1	101.1	101.4	100.8	100.1	100.2	100.4	100.5
Fo	95.99	96.44	96.70	94.69	94.93	95.72	95.25	96.02	96.35	95.15	94.96
Fa	1.66	1.56	1.57	2.56	1.96	2.53	2.15	1.97	1.28	2.46	2.78

Abbreviations: Px: pyroxene; Ol: Olivine; Wo: Wollastonite; En: Enstatite; Fs: Ferrosilite; Fo: Forsterite; Fa: Fayalite

Online Resource Table OR2

Trace element concentrations of drilled micro-samples of experimental glass.

Table 2

PC448-V9 (60 s)				PC441-V1 (90 s)			PC442-V4 (60 s)			PC441-V2 (90 s)			
Experiment location	L-V9.1	L-V9.3	L-V9.2	L-V1.1	L-V1.2	L-V1.3	D-V4.1	D-V4.2	D-V4.3	D-V2.1	D-V2.2	D-V2.3	D-V2.4
Ti ⁴⁹	0.5	0.8	0.6	0.6	0.8	0.3	0.7	0.8	0.9	0.8	0.8	0.9	1.1
Rb ⁸⁵	73.8	122.3	102.7	41.8	155.6	64.2	178.2	170.8	217.7	139.4	121.6	122.5	114.0
Sr ⁸⁸	423.0	605.9	482.3	395.6	596.3	227.7	477.7	534.1	578.1	547.2	563.3	547.8	601.6
Y ⁸⁹	11.4	18.2	19.4	3.2	19.9	7.4	17.4	18.6	23.0	15.6	17.6	19.6	22.1
Zr ⁹⁰	78.0	131.9	101.8	73.4	136.4	51.2	116.6	131.2	143.6	132.3	134.3	133.0	120.3
Nb ⁹³	4.7	17.1	11.7	5.4	12.0	6.2	14.2	15.8	12.8	14.3	4.8	7.2	9.7
Ba ¹³⁷	560.2	952.9	712.7	459.5	1107.0	388.9	872.9	899.8	1049.5	932.9	956.3	939.0	941.4
La ¹³⁹	13.6	22.5	41.9	3.6	23.5	8.9	20.7	24.6	35.5	20.4	22.5	27.2	36.1
Ce ¹⁴⁰	28.6	46.3	43.6	7.5	49.2	18.0	42.9	50.2	58.8	45.8	48.7	49.9	66.3
Pr ¹⁴¹	3.8	6.4	10.3	1.0	6.6	2.5	5.9	6.8	9.4	5.8	6.2	7.2	9.7
Nd ¹⁴³	15.7	25.7	40.5	4.2	27.8	10.4	24.2	27.3	36.6	24.0	25.3	29.3	38.6
Sm ¹⁴⁷	3.4	5.5	6.7	1.0	5.9	2.1	4.9	5.6	7.5	4.9	5.2	6.1	7.3
Sm ¹⁴⁹	3.6	5.5	7.3	0.9	5.7	2.3	4.9	5.8	7.4	5.0	5.4	6.0	7.7
Eu ¹⁵¹	1.0	1.6	1.8	0.3	1.6	0.6	1.5	1.6	1.9	1.4	1.5	1.6	2.1
Gd ¹⁵⁷	3.4	5.7	7.1	0.9	5.8	2.2	5.0	5.7	6.7	4.7	5.3	5.7	7.2
Dy ¹⁶¹	2.2	3.3	3.4	0.6	3.8	1.4	3.3	3.4	4.2	2.9	3.3	3.6	4.1
Er ¹⁶⁶	1.0	1.6	1.6	0.3	1.6	0.6	1.4	1.6	2.0	1.3	1.5	1.7	1.9
Yb ¹⁷²	0.9	1.5	1.4	0.2	1.5	0.5	1.4	1.4	1.6	1.2	1.4	1.5	1.7

Table 2 (continued)

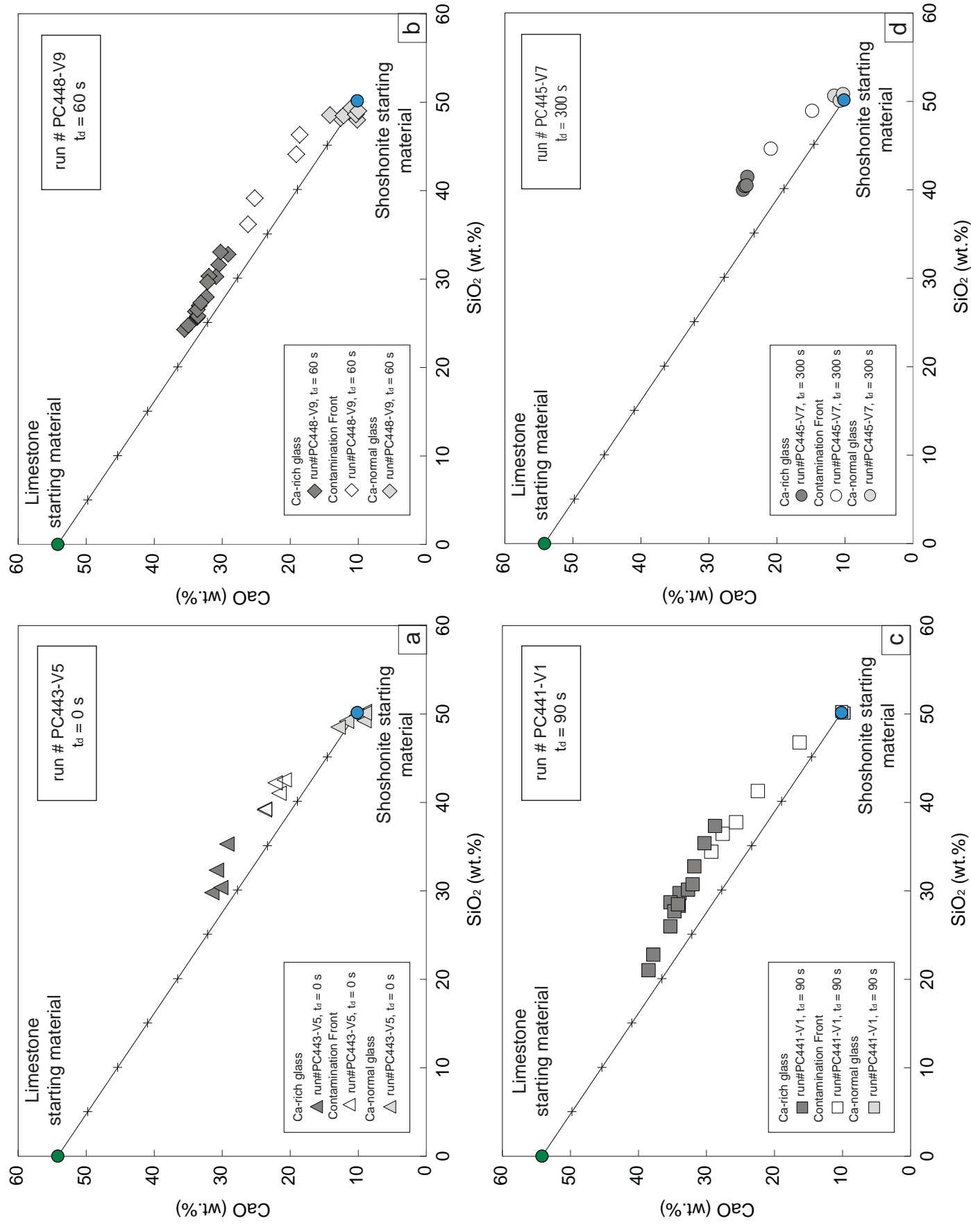
PC448-V9 (60 s)				PC441-V1 (90 s)			PC442-V4 (60 s)			PC441-V2 (90 s)			
Experiment, location	L-V9.1	L-V9.3	L-V9.2	L-V1.1	L-V1.2	L-V1.3	D-V4.1	D-V4.2	D-V4.3	D-V2.1	D-V2.2	D-V2.3	D-V2.4
Lu ¹⁷⁵	0.10	0.16	0.16	0.03	0.18	0.06	0.14	0.15	0.19	0.13	0.15	0.17	0.19
Pb ²⁰⁸	38.98	24.75	18.67	13.48	23.96	7.88	36.85	21.5	38.11	18.71	20.2	20.1	20.46
Th ²³²	4.81	7.69	12.84	1.30	8.13	3.11	7.30	8.08	10.65	6.33	7.22	8.83	10.20
U ²³⁸	2.41	3.22	2.61	2.53	3.07	1.23	2.71	3.13	3.21	2.94	3.30	3.04	2.68
Tb ¹⁵⁹	0.01	0.02	0.02	0.00	0.02	0.01	0.02	0.02	0.02	0.01	0.02	0.02	0.02

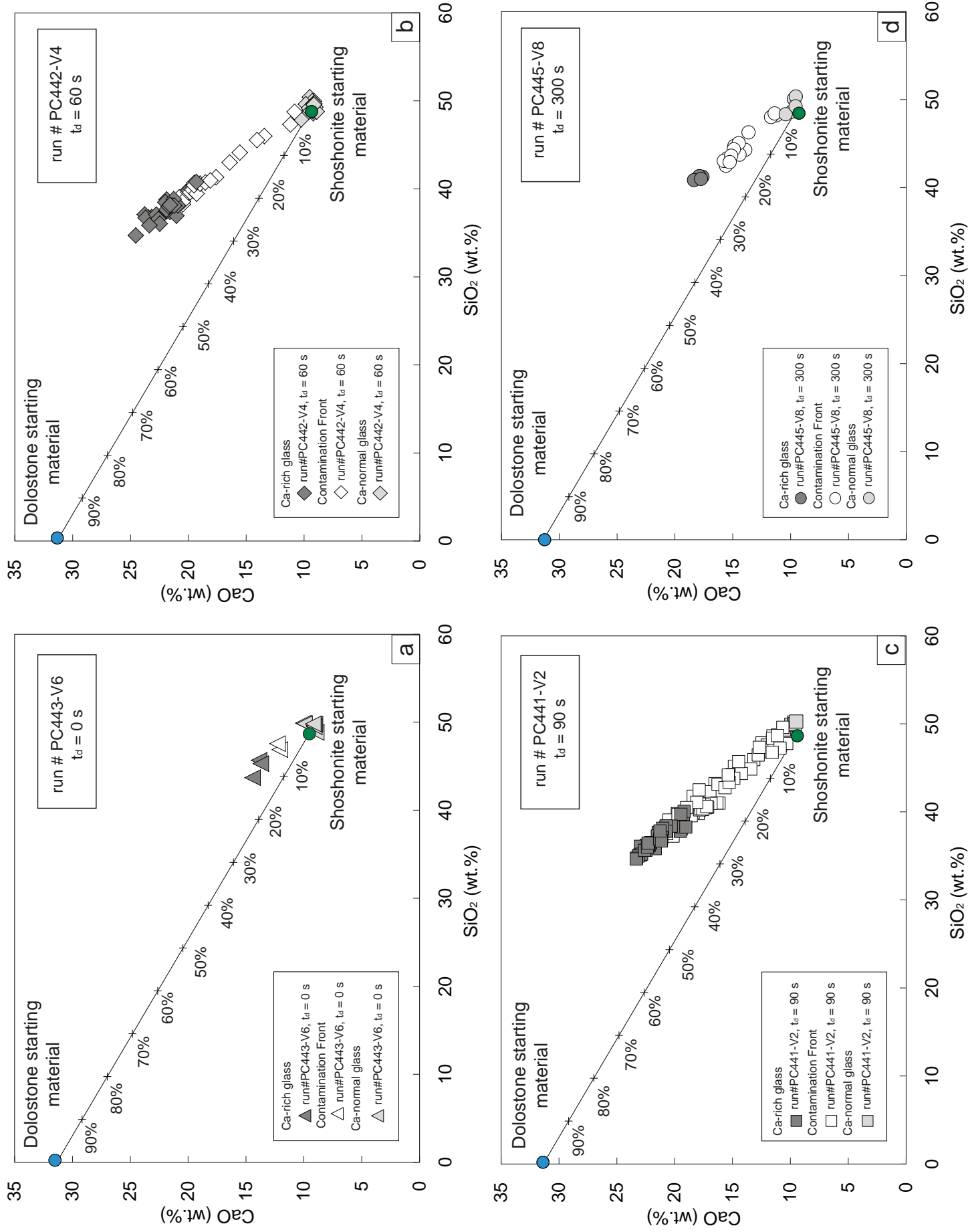
Concentrations are given in ppm. Location numbers refer to sampled areas in the experiments, as shown in **Fig.5**.

Online Resource 3

Figure OR1. Mixing trends between starting materials in run durations of: 0 s (**a**), 60 s (**b**), 90 s (**c**) and 300 s (**d**) for limestone-bearing experiments. For $t_d = 0$ and 60 s, the system has locally assimilated up to 40% and 50% carbonate-derived CaO respectively. For $t_d = 90$ s the system reached its maximum of mixing, being able to assimilate ~ 60% of carbonate-derived CaO. For $t_d = 300$ s, the system has almost homogenised giving an intermediate mixing ratio overall.

Figure OR2. Mixing trends between starting materials in runs at: 0 s (**a**), 60 s (**b**), 90 s (**c**) and 300 s (**d**) for dolostone-bearing experiments. For $t_d = 0$ s, around 10 - 15% of carbonate-derived CaO has been assimilated. For $t_d = 60$ and 90 s, the system reached its maximum degree of mixing of ~ 25-40% of carbonate-derived CaO. For $t_d = 300$ s, ~ 25% of the carbonate-derived CaO is assimilated in total due to an almost homogenous melt.





Online Resource 4

Apparent diffusion coefficients (D) were calculated from the compositional profiles through the limestone-bearing experiments by applying the expression

$$D = x^2/t$$

where D is the apparent diffusion coefficient in m^2/s , x is the location of a specific element along the traverse in meters and t is the experimental time in seconds. This allows us to evaluate the effectiveness of diffusion of an element over a distance across the contamination front and estimate the efficiency of diffusion relative to other processes. The results of these calculations are summarised in **Table OR1**.

Table OR1: *Measured diffusion coefficients*

Limestones							
run#	T (°C)	time (s)	Traverses	D (m^2/s)			
				SiO ₂	Al ₂ O ₃	MgO	CaO
PC442-V3	1200	60	1	$8.6 \cdot 10^{-7}$	$4.0 \cdot 10^{-10}$	$1.1 \cdot 10^{-8}$	$7.5 \cdot 10^{-9}$
PC448-V9	1200	60	1	$2.0 \cdot 10^{-11}$	$2.0 \cdot 10^{-11}$	$1.8 \cdot 10^{-10}$	$6.3 \cdot 10^{-6}$
			2	$3.9 \cdot 10^{-9}$	$3.9 \cdot 10^{-9}$	$3.9 \cdot 10^{-9}$	$2.0 \cdot 10^{-11}$
PC441-V1	1200	90	1	$2.5 \cdot 10^{-9}$	$1.9 \cdot 10^{-9}$	$2.5 \cdot 10^{-9}$	$4.5 \cdot 10^{-10}$
PC432-V7	1200	300	1	$8.5 \cdot 10^{-10}$	$8.5 \cdot 10^{-10}$	$8.5 \cdot 10^{-10}$	$3.4 \cdot 10^{-11}$